Numerical methods for phase retrieval

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Numerical methods for phase retrieval

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In this work we consider the problem of reconstruction of a signal from the magnitude of its Fourier transform, also known as phase retrieval. The problem arises in many areas of astronomy, crystallography, optics, and coherent diffraction imaging (CDI). Our main goal is to develop an efficient reconstruction method based on continuous optimization techniques. Unlike current reconstruction methods, which are based on alternating projections, our approach leads to a much faster and more robust method. However, all previous attempts to employ continuous optimization methods, such as Newton-type algorithms, to the phase retrieval problem failed. In this work we provide an explanation for this failure, and based on this explanation we devise a sufficient condition that allows development of new reconstruction methods—approximately known Fourier phase. We demonstrate that a rough (up to $\pi/2$ radians) Fourier phase estimate practically guarantees successful reconstruction by any reasonable method. We also present a new reconstruction method whose reconstruction time is orders of magnitude faster than that of the current method-of-choice in phase retrieval—Hybrid Input-Output (HIO). Moreover, our method is capable of successful reconstruction even in the situations where HIO is known to fail. We also extended our method to other applications: Fourier domain holography, and interferometry.

Additionally we developed a new sparsity-based method for sub-wavelength CDI. Using this method we demonstrated experimental resolution exceeding several times the physical limit imposed by the diffraction light properties (so called diffraction limit).

*The work on sub-wavelength CDI was done in collaboration with Prof. M. Segev’s group from the Technion Physics Department, Solid State Institute.
1 Introduction

1.1 Motivation

Recent development of nanotechnology has resulted in great interest in imaging techniques suitable for visualization of nano-structures. One of the most promising techniques for such high resolution imaging is Coherent Diffraction Imaging (CDI). In CDI, a highly coherent beam of X-rays or electrons is incident on a specimen, generating a diffraction pattern. Under certain conditions the diffracted wavefront is approximately equal (within a scale factor) to the Fourier transform of the specimen. After being recorded by a CCD sensor, the diffraction pattern is used to reconstruct the specimen (Sayre, 1952; Miao et al., 1999; Quiney, 2010). Effectively, in CDI we replace the objective lens of a typical microscope with a software algorithm. The advantage in using no lenses is that the final image is aberration-free and the final resolution is only diffraction and dose limited, that is, dependent only on the wavelength, aperture size and exposure time. This process is illustrated in Figure 1.1.

![Fig 1.1: CDI process](image)

The method has been successfully applied to visualizing a variety of nano-structures, such as carbon nano-tubes (Zuo et al., 2003), defects inside nano-crystals (Pfeifer et al., 2006), proteins, and more (Neutze et al., 2000; Chapman et al., 2006; Chapman et al., 2007). Furthermore, exactly the same problem—the reconstruction of a signal from the magnitude of its Fourier transform—arises in may other areas of science. Notable examples include astronomy, crystallography, and speckle interferometry.
It is important to note that, due to the physical nature of the sensor, we are limited to recording only the intensity (squared amplitude) of the diffracted wave, hence its phase is lost. As will be shown later, this loss of the phase leaves us with highly incomplete data, which makes the problem of reconstruction hard.

1.2 Data acquisition model

Of course, in the real world, the sought object $x(t)$ and its Fourier transform (denoted by $\hat{x}(\omega)$) are both continuous functions of $t$ and $w$, respectively, where $t$ and $w$ are multidimensional coordinate vectors. Furthermore, the support of $x(t)$ is limited, which means that $\hat{x}(\omega)$ is spread over all frequencies: from $-\infty$ to $\infty$. However, during the data acquisition process we capture only a finite extent in the Fourier domain and all further processing is done on digital computers. This naturally leads to discrete approximations of $x(t)$ and $\hat{x}(\omega)$, that are well justified in view of the finite resolution that stems from the measurements and, in general, from the fact that all optical systems have resolution limits. Given that $x[n]$ (an adequate sampling of $x(t)$) contains $N$ points (for simplicity we assume that $x$ is one-dimensional—generalization for the multi-dimensional case is straightforward) we assume that $x[n]$ vanishes outside the interval $[0, N-1]$. Furthermore, if we assume, without loss of generality, that the physical extent of $x$ is unity we immediately conclude that the sampling rate in the Fourier domain must be $1/N$ to acquire the measurements that are related to $x[n]$ via the Discrete Fourier Transform (DFT). However, a more thorough examination of the problem yields a higher sampling rate requirement. Recall that we record only the intensity of the diffraction pattern. This intensity can be represented as follows:

$$I(\omega) = |\hat{x}(\omega)|^2 = \overline{\hat{x}(\omega)} \circ \hat{x}(\omega), \quad (1.1)$$

where the overbar denotes the complex conjugate and $\circ$ denotes element-wise multiplication. Hence, the inverse Fourier transform of the measured intensity $I(\omega)$ results in the auto-correlation function (denoted by $\star$) of the sought object,

$$\mathcal{F}^{-1}[I(\omega)] = x(t) \star x(t). \quad (1.2)$$

Obviously, the autocorrelation $x(t) \star x(t)$ has support that is twice as large as the support of $x(t)$ (in each dimension), therefore, the diffraction pattern intensity must be sampled with the rate two times higher than $1/N$ to capture all the information about the auto-correlation function. To this end, we always assume that the signal $x(t)$ (or $x[n]$) is “padded” with zeros so that its size is doubled (in each dimension). This requirement is an approximation to the physical constraint on $x(t)$ having finite support. Without adding it into the reconstruction scheme, the problem would be severely undetermined with multiple solutions that are unrelated to the original signal $x$. To illustrate the last claim, one can imagine the case where $x[n]$ is reconstructed from its Fourier magnitude without additional constraints. Obviously, any choice of
the Fourier phase will give rise to a valid solution which, unfortunately, has little to do with the sought signal.

1.3 Reconstruction from incomplete Fourier data

Before we proceed to the main subject of this work—the reconstruction of a signal from the magnitude of its Fourier transform—let us consider a number of toy problems, where we evaluate the importance of different parts of the Fourier transform.

The Fourier transform is, in general, complex. There are two common representations of a complex number: one is the sum of its real (Re) and imaginary (Im) parts

\[ z = \text{Re} + j \text{Im}, \]  

(1.3)

and the other is the product of its magnitude (r) and the complex exponent of its phase \( e^{j\phi} \),

\[ z = r e^{j\phi}. \]  

(1.4)

From these formulas it is not clear whether one part of the representation is more important than the other. Below we demonstrate that the real and the imaginary parts carry about equal amount of information and the loss of one of them can often be recovered. However, the phase carries most of the information, and consequently, its loss is more difficult to overcome.

1.3.1 Reconstruction from the real part

Let us assume that \( x[n] \) is a real one-dimensional signal\(^\ast\). Furthermore, we assume that \( x \) vanishes outside the interval \([0, N - 1]\), specifically, we assume that \( x[n] = 0 \) for \( n = -1, -2, \ldots, -(N - 1) \). Recall that any real signal can be represented uniquely as a sum of two signals

\[ x = x_e + x_o, \]  

(1.5)

where \( x_e \) is even and \( x_o \) is odd\(^\dagger\) (see Figure 1.2). It can be easily shown that

\[ x_e = \frac{x[n] + x[-n]}{2}, \quad x_o = \frac{x[n] - x[-n]}{2}. \]  

(1.6)

Recall also that the Fourier transform of an even signal is real and that of an odd signal is purely imaginary. Hence, we conclude that the real part of \( \hat{x} \) is nothing but the Fourier transform of \( x_e \). Thus, we can obtain the even part \( x_e \) by the inverse Fourier transform of the real part of \( \hat{x} \). Furthermore, reconstructing \( x \) from \( x_e \) is trivial—we should take the right-hand side \((n \geq 0)\) of \( x_e \) multiplied by two everywhere except the origin (see Figure 1.2).

\(^\ast\)A generalization to a complex multidimensional \( x \) is straightforward.

\(^\dagger\)In the complex case, \( x_e \) is Hermitian, and \( x_o \) is anti-Hermitian.
Figure 1.2: Signal decomposition into even and odd parts: (a) original signal $x[n]$, (b) its reversed version $x[-n]$, (c) the even part, and (d) the odd part.
1.3.2 Reconstruction from the imaginary part

The reconstruction from the imaginary part is also easy. The method is very similar to the reconstruction from the real part. The only difference is that now we obtain the odd part of the sought signal. But, again, taking the right-hand side of \( x_o \) and multiplying it by two we obtain the original signal \( x \) everywhere besides the origin.

For the most general case, where \( x \) is complex, it is easy to show that the missing imaginary part leads to almost perfect reconstruction—only \( \text{Im}(x[0]) \) is lost. Similarly, when reconstructing from the imaginary part of \( \hat{x} \)—only \( \text{Re}(x[n]) \) is lost. Therefore, we can conclude that the real and the imaginary parts of the Fourier transform carry the same amount of information and losing either one of them can be easily overcome if \( x[n] \) is sufficiently padded with zeros and vanishes for \( n < 0 \).

1.3.3 Reconstruction from the phase

Now we switch to the second (polar) representation of complex numbers and consider first the situation where the magnitude is lost, namely, reconstruction from the Fourier phase. Several researchers (see (Hayes, 1982; Hayes et al., 1980; Millane, 1996; Millane, 1990; Oppenheim & Lim, 1981)) showed that sufficiently padded signals can be reconstructed from the Fourier phase. However, the reconstruction is possible only up to a scale factor, that is, one obtains \( \alpha x \) for some real positive scalar \( \alpha \). Furthermore, it was shown that this is the only type of non-uniqueness possible in signal reconstruction from the Fourier phase (Hayes et al., 1980).

It seems that all previous works used a variation of the method of alternating projections (see Section 3.1). However, the problem can easily be represented as linear programming, for which much more efficient algorithms exist. Actually, we solve a variation of this problem, where the Fourier phase is known to lie within certain interval in Chapters 5 and 6, and our method is much faster than the method of alternating projections.

1.3.4 Reconstruction from the magnitude

We finally arrive at the case that is the main subject of this research—reconstruction from the Fourier magnitude alone. It turns out that this case of incomplete Fourier information is the most difficult among the four possibilities considered here. This phenomenon can be related to the fact that the Fourier phase carries the majority of the information in a signal. As an informal “proof” of the latter claim, look at Figure 1.3, where we exchange the Fourier phase between two different images, while keeping the Fourier magnitude intact. The results show, unmistakably, an exchange of the images.

It turns out that this problem is very different in the one-dimensional case and in the multi-dimensional case. The former suffers from multiplicity of possible solutions (no matter how much padding we use in \( x \)), while the latter is usually less prone to multiple solutions (aside from trivial transformations: lateral shifts, axis reversal,
and constant phase factor), but finding a solution is very difficult. An explanation for this phenomenon is given in Chapter 2.
Figure 1.3: The importance of phase in signals—given two images (a) a dog, and (b) a cat, we exchange their Fourier phase while keeping the magnitude. The results, unmistakably, show the cat instead of the dog (c), and the dog instead of the cat (d).
1.4 A short overview

After this short introduction we are ready to proceed to the main part of this work. In the next chapter we present the mathematical foundations of the phase retrieval problem: our main concern is uniqueness of the reconstruction, moreover, we demonstrate why the one-dimensional and the multi-dimensional cases behave extremely differently. In Chapter 3 we present the methods that are used today for phase retrieval. These chapters are based on a compilation of known facts and do not contain our original research. Our development starts in Chapter 4, where we introduce machinery for efficient optimization of a real function of complex argument. Moreover, we find and analyze the eigen-decomposition of the Hessian of one of the most frequently used objective function in phase retrieval. Chapters 5 and 6 are dedicated to a variant of the phase retrieval problem where the Fourier phase is not lost completely, but rather a rough estimate of it is available. We demonstrate empirically and then prove mathematically that this scenario leads to a new family of algorithms that are orders of magnitude faster than the existing methods. The ideas from these chapters are taken into the new field of the Fourier domain holography in Chapters 7, and 8 where we develop a new method of reconstruction and present guide rules for a reference beam design that lead to fast and robust reconstruction. A new type of prior knowledge—image sparsity—is exploited in Chapter 9 where we present a CDI with resolution several times higher than the maximal theoretical limit. Finally, in Chapter 10 we present concluding remarks and provide reference to the works that was done in the course of this Ph.D. research but have not been included in this thesis. For example, we used Fienup’s Hybrid Input-Output algorithm (see Section 3.2) for creation of Grassmanian matrices (Osherovich et al., 2009a)—an interesting work that was left outside because it is not related to the phase retrieval problem.
# 2 Mathematical foundations

In this chapter we review some mathematical properties of the phase retrieval problem. Our main concern is the uniqueness properties of the solution, which turns to be different in the one-dimensional and multi-dimensional cases.

## 2.1 One-dimensional signals

### 2.1.1 Continuous-time case

What can be said about two signals \( g_1(t) \) and \( g_2(t) \) having the same power spectrum? Obviously,

\[
|\hat{g}_1(\omega)|^2 = |\hat{g}_2(\omega)|^2 \tag{2.1}
\]

means that

\[
|\hat{g}_1(\omega)| = |\hat{g}_2(\omega)|. \tag{2.2}
\]

This, in turn, leads to the following relation

\[
\hat{g}_2(\omega) = \hat{g}_1(\omega)e^{j\phi(\omega)}. \tag{2.3}
\]

for some real-valued function \( \phi(\omega) \). Hence, if \( g_2(t) \) has the same power spectrum as \( g_1(t) \), then \( g_2(t) \) must be a result of passing \( g_1(t) \) through an all-pass filter. Moreover, it means that there are infinitely many (an uncountable set of) functions having the same power spectrum, because any choice of \( \phi(\omega) \) will lead to some function

\[
g_2(t) = \frac{1}{2\pi} \int_{-\infty}^{\infty} \hat{g}_1(\omega)e^{j\phi(\omega)}e^{j\omega t} \, d\omega. \tag{2.4}
\]

This observation, however, is not particularly helpful as we are not interested in arbitrary signals. Our interest is limited to signals that are physically feasible. Such signals, for example, must have finite support and finite energy. That is, we assume that

\[
g(t) = 0, \text{ for } |t| > \frac{T}{2}, \tag{2.5}
\]

and

\[
\int_{-T/2}^{T/2} |g(t)|^2 \, d(t) < \infty. \tag{2.6}
\]
Under these restrictions it is not clear anymore that there still exist infinitely many signals having the same power spectrum. Nevertheless, it was shown that even under these conditions the number of signals having the same power spectrum can be infinite. A fairly complete treatment of this problem is given by Hofstetter (1964) who considered the problem from a different point of view: finding of all possible time-limited signals having the same autocorrelation function as a given signal. However, it is exactly the same problem due to the direct connection between the autocorrelation and the power spectrum. Although Hofstetter’s work was done for continuous-time (analogue) signals and not for discrete signals that we encounter in computer algorithms, we believe it is more instructive to start with this case and proceed then to the discrete version of the problem. Hence, we present next the main results and derivations found in (Hofstetter, 1964).

Under the conditions (2.5) and (2.6), \( g(t) \) is absolutely integrable
\[
\int_{-T/2}^{T/2} |g(t)| \, dt < \infty. \tag{2.7}
\]
and its Laplace transform,
\[
G(s) \equiv \mathcal{L}[g(t)] = \int_{-T/2}^{T/2} g(t) e^{-st} \, dt, \tag{2.8}
\]
converges for all complex \( s = \sigma + j\omega \). The function \( G(s) \) is thus analytic in the entire \( s \) plane and, because of Equation (2.6), square integrable. The signal \( g(t) \) can always be recovered from \( G(s) \) by means of the inverse transform
\[
g(t) = \frac{1}{2\pi} \int_{-\infty}^{\infty} G(j\omega) e^{j\omega t} \, d\omega. \tag{2.9}
\]
From the definition of the Laplace transform, it is obvious that it is a generalization of the Fourier transform, and the relation between the two is very simple:
\[
\hat{g}(\omega) = \mathcal{F}[g(t)] = \mathcal{L}[g(t)] \bigg|_{s=j\omega} = G(j\omega). \tag{2.10}
\]

With these preliminaries in hand we proceed to the autocorrelation function of \( g(t) \) defined as
\[
r(\tau) \equiv g(t) \star g(t) = \int_{-T/2}^{T/2} \bar{g}(t) g(t+\tau) \, d\tau. \tag{2.11}
\]
It can be readily shown that \( r(\tau) \) also has finite support: it vanishes outside the interval \([-T, T]\). Moreover, similarly to \( g(t) \), \( r(\tau) \) is square and absolutely integrable.

\*Here we do not count the trivial multiplication by a phase factor.
Therefore, its Laplace transform,

\[ R(s) = \int_{-T}^{T} r(\tau)e^{-s\tau} d\tau , \] (2.12)

exists, and is analytic in the entire \( s \) plane. Of course, it and can be used to recover \( r(\tau) \) by the means of the inverse transform

\[ r(\tau) = \frac{1}{2\pi} \int_{-\infty}^{\infty} R(j\omega)e^{j\omega\tau} d\omega . \] (2.13)

Since our main goal is to design signals with the same autocorrelation function, we need to find the relation between the Laplace transform of a signal and the Laplace transform of its autocorrelation function. The development is straightforward

\[ R(s) = \int_{-\infty}^{\infty} (g(t) \ast g(t)) e^{-s\tau} d\tau \] (2.14)
\[ = \int_{-\infty}^{\infty} \left[ \int_{-\infty}^{\infty} \bar{g}(t)g(t+\tau) dt \right] e^{-s\tau} d\tau \] (2.15)
\[ = \int_{-\infty}^{\infty} \bar{g}(t)e^{st} dt \int_{-\infty}^{\infty} g(t+\tau)e^{-s(t+\tau)} d\tau \] (2.16)
\[ = \overline{G(-s)}G(s) . \] (2.17)

In the transition from (2.16) to (2.17) we used the fact that

\[ L[\bar{g}(t)] = \overline{G(s)} . \] (2.18)

Hence, if two signals \( g_1(t) \), and \( g_2(t) \) have the same autocorrelation function they must obey the following equality for all \( s \):

\[ G_1(-\bar{s})G_1(s) = \overline{G_2(-\bar{s})G_2(s)} . \] (2.19)

This includes \( s = j\omega \), which gives the equality between their power spectra

\[ \overline{G_1(-j\omega)}G_1(j\omega) = \overline{G_2(-j\omega)}G_2(j\omega) \]
\[ \overline{G_1(j\omega)G_1(j\omega)} = \overline{G_2(j\omega)G_2(j\omega)} \]
\[ |G_1(j\omega)|^2 = |G_2(j\omega)|^2 , \] (2.20)

as expected.

The necessity to introduce the Laplace transform will become clear after we present a family of all-pass filters that preserve the support bounds of \( g(t) \). It is shown in (Hofstetter, 1964) that if \( s_0 \) is a zero of \( G_1(s) \), that is, if

\[ G_1(s_0) = \int_{-\infty}^{\infty} g_1(t)e^{-s_0t} dt = 0 , \] (2.21)
then the all-pass filter \( h(t) \) whose transfer function \( H(s) \) is given by

\[
H(s) = \frac{s + \bar{s}_0}{s - s_0}
\]  

(2.22)

will not spread a given signal \( g_1(t) \) outside the original interval \([-T/2, T/2]\). It is easy to verify that

\[
|H(j\omega)|^2 = H(j\omega)H(j\omega) = \left( \frac{j\omega + \bar{s}_0}{j\omega - s_0} \right) \left( \frac{-j\omega + s_0}{-j\omega - \bar{s}_0} \right) = 1.
\]  

(2.23)

Hence, \( h(t) \) is indeed an all-pass filter because \( |H(j\omega)| = 1 \). To demonstrate that \( h(t) \) preserves the support, let us find its explicit representation. To this end, it is more convenient to split it into the following two filters

\[
H(s) = 1 + \frac{s_0 + \bar{s}_0}{s - s_0}.
\]  

(2.24)

In this form, it is easy to see that the impulse response \( h(t) \) that corresponds to \( H(s) \) is given by

\[
h(t) = \frac{1}{2\pi} \int_{-\infty}^{\infty} \left( 1 + \frac{s_0 + \bar{s}_0}{j\omega - s_0} \right) e^{j\omega t} d\omega
\]  

(2.25)

\[
= \delta(t) + \frac{s_0 + \bar{s}_0}{2\pi} \int_{-\infty}^{\infty} \frac{e^{j\omega t} e^{-s_0 t}}{j\omega - s_0} d\omega
\]  

(2.26)

\[
= \delta(t) + (s_0 + \bar{s}_0) e^{s_0 t} U(t),
\]  

(2.27)

where \( U(t) \) denotes the Heaviside step function. Hence, if a time-limited input \( g_1(t) \) were provided to the system described by \( H(s) \), the output \( g_2(t) \) would be \( g_1(t) \) plus the convolution of \( g_1(t) \) with the filter \( h_1(t) = (s_0 + \bar{s}_0) e^{s_0 t} U(t) \). To show that the output \( g_2(t) \) vanishes outside the interval \([-T/2, T/2]\) we must show that the latter term (convolution) vanishes outside this interval. The proof is straightforward:

\[
g_{21}(t) = g_1(t) \otimes h_1(t)
\]  

(2.28)

\[
ge_{T/2} -T/2 \int g_1(\tau)h_1(t - \tau) d\tau
\]  

(2.29)

\[
= \begin{cases} 
0 & \text{if } t < -\frac{T}{2}, \\
(s_0 + \bar{s}_0) \int_{-T/2}^{t} g_1(\tau) e^{s_0 (t-\tau)} d\tau & \text{if } -\frac{T}{2} \leq t \leq \frac{T}{2}, \\
(s_0 + \bar{s}_0) \int_{-T/2}^{T/2} g_1(\tau) e^{s_0 (t-\tau)} d\tau & \text{if } t > \frac{T}{2},
\end{cases}
\]  

(2.30)

where \( \otimes \) denotes convolution. Note that \( g_{21}(t) \) vanishes for \( t > T/2 \), because the last
line in the above equation reads

\[
(s_0 + \bar{s}_0) \int_{-T/2}^{T/2} g_1(\tau) e^{s_0(t-\tau)} d\tau = (s_0 + \bar{s}_0)e^{s_0t} \int_{-T/2}^{T/2} g_1(\tau)e^{s_0\tau} d\tau
\]

\[
= (s_0 + \bar{s}_0)e^{s_0t}G_1(s_0),
\]

(2.31)

and \( s_0 \) is a root of \( G_1(s) \). Hence, the support of \( g_2 \) is not wider than the support of \( g_1 \), that is, it vanishes outside the interval \([-T/2, T/2]\). It is important to emphasize the way \( G_2(s) \) (equivalently, \( g_2(t) \)) is obtained from \( G_1(s) \) (equivalently, \( g_1(t) \)):

\[
G_2(s) = G_1(s) \frac{s + \bar{s}_0}{s - s_0},
\]

(2.32)

where \( s_0 \) is a root of \( G_1(s) \). This formula gives an obvious way to obtain a new time-limited signal \( g_2(t) \) from a given time-limited signal \( g_1(t) \) such that the autocorrelation of the two is equal. The algorithm is really simple, indeed:

1. Laplace transform \( g_1(t) \) to obtain \( G_1(s) \)

2. Choose \( k \) non-zero roots of \( G_1(s) \): \( \{s_i\}_{i=1}^{k} \) and replace them with their negative conjugates, to obtain \( G_2(s) \)

\[
G_2(s) = G_1(s) \prod_{i=1}^{k} \frac{s + \bar{s}_i}{s - s_i}
\]

(2.33)

3. Inverse Laplace transform \( G_2(s) \) to obtain \( g_2(s) \)

The fact that the new signal \( g_2(t) \) vanishes outside the interval \([-T/2, T/2]\) was proven above. The fact that the two signals have the same autocorrelation function and, hence, the same power spectrum is readily obtained by simple calculations:

\[
\mathcal{L}[g_2(t) \ast g_2(t)] = \frac{G_2(-\bar{s})G_2(s)}{G_1(-s) \prod_{i=1}^{k} \frac{s + \bar{s}_i}{s - s_i}} G_1(s) \prod_{i=1}^{k} \frac{s + \bar{s}_i}{s - s_i}
\]

\[
= \frac{G_1(-\bar{s})G_1(s)}{G_1(-s) \prod_{i=1}^{k} \frac{s - s_i}{s - \bar{s}_i}} G_1(s) \prod_{i=1}^{k} \frac{s + \bar{s}_i}{s - s_i}
\]

\[
= \frac{G_1(-\bar{s})G_1(s)}{G_1(-s) \prod_{i=1}^{k} \frac{s - s_i}{s + \bar{s}_i}} G_1(s) \prod_{i=1}^{k} \frac{s + \bar{s}_i}{s - s_i}
\]

(2.34)

This, actually, leads us to an algorithm for generating new time-limited signals, all having the same autocorrelation function. The algorithm simply replaces one or
more non-zero roots of the Laplace transform of a given signal with their negative conjugates. Moreover, it can be proven that this approach can generate all possible signals with provided time support and autocorrelation function. A rigorous proof of the last statement actually expands the current result to an infinite set of roots. The proof is not difficult but we do not present it here. The interested reader can find it in (Hofstetter, 1964). For our discussion it is more important to note that a set of $k$ non-zero roots of $G_1(s)$ gives rise to $2^k$ new signals with the same time support and autocorrelation function. Of course, depending on additional constraints, some of these signals may not be feasible. For example, if our attention is restricted to real-valued signals, the zeros of $G_1(s)$ must occur in conjugate pairs. Hence, to generate a new real-valued signal we must replace the corresponding pair and not a single root.

Below we present a number of examples of this technique (from (Hofstetter, 1964))

**Example 1** Assume that $a < 0$ and let $g_1(t)$ be given by

$$g_1(t) = \begin{cases} 0 & \text{if } |t| > 1, \\ -e^{at} & \text{if } -1 \leq t < 0, \\ e^{at} & \text{if } 0 < t \leq 1. \end{cases} \quad (2.35)$$

Calculating its Laplace transform gives us

$$G_1(s) = \int_{-1}^{0} -e^{(a-s)t} \, dt + \int_{0}^{1} e^{(a-s)t} \, dt$$

$$= 2 \frac{1 - \cosh(s-a)}{s-a}. \quad (2.36)$$

Since $G_1(a) = 0$, we can create a new signal $g_2(t)$ by passing $g_1(t)$ through the all-pass filter given by

$$H(s) = \frac{s+a}{s-a}. \quad (2.37)$$

The impulse response of this filter is

$$h(t) = \delta(t) + 2aU(t)e^{at}. \quad (2.38)$$

Hence, the output of the filter is

$$g_2(t) = \begin{cases} 0 & \text{if } |t| > 1, \\ -e^{at}(2at + 2a + 1) & \text{if } -1 \leq t < 0, \\ e^{at}(2at - 2a + 1) & \text{if } 0 < t \leq 1. \end{cases} \quad (2.39)$$

Figure 2.1 below depicts the two signals $g_1(t)$, $g_2(t)$, and their common autocorrelation function.
Figure 2.1: Example 1: two pulses (a), and (b), and their common autocorrelation function (c).

Example 2 Let $g_1(t)$ be given by

\[
g_1(t) = \begin{cases} 
0 & \text{if } |t| > 1, \\
e^{at} & \text{if } |t| \leq 1.
\end{cases}
\]  

Then

\[
G_1(t) = \int_{-1}^{1} e^{(a-s)t} \, dt = 2 \frac{\sinh(s - a)}{s - a}.
\]  

The zeros of $G_1$ are located at the points $s_k$ given by

\[
s_k = a + jk\pi, \quad k = \pm 1, \pm 2.
\]

We can construct a real-valued signal having the same support and autocorre-
lation function as $g_1(t)$ by letting

$$G_2(s) = G_1(s)H(s),$$  \hfill (2.43)

where

$$H(s) = \frac{s + a - j\pi \cdot s + a - j\pi}{s - a - j\pi \cdot s - a + j\pi} = \frac{(s + a)^2 + \pi^2}{(s - a)^2 + \pi^2},$$  \hfill (2.44)

The impulse response $h(t)$ of this all-pass filter is easily found to be

$$h(t) = \delta(t) + 4aU(t)e^{at}\left(\cos(\pi t) + \frac{a}{\pi}\sin(\pi t)\right),$$  \hfill (2.45)

and the convolution between $g_1(t)$ and $h(t)$ yields the result

$$g_2(t) = \begin{cases} 
0 & \text{if } |t| > 1, \\
\frac{4a}{\pi} e^{at}\left(\frac{a}{\pi}\cos(\pi t) - \sin(\pi t) + \frac{a}{\pi}\right) + e^{at} & \text{if } |t| \leq 1. 
\end{cases}$$  \hfill (2.46)

Figure 2.2 depicts the signals $g_1(t)$, $g_2(t)$; and their common autocorrelation function.
2.1.2 Discrete-time case

The derivations in the previous section describe completely the theory of continuous-time one-dimensional signals. However, for algorithms that run on a digital computer, we need a similar theory for discrete-time signals, that is, for $g(t)$ that is specified on a finite set of points

$$g(t) = \sum_n g_n \delta(t - t_n),$$  \hspace{1cm} (2.47)

where $t_n = n\Delta$. The corresponding Fourier transform is, of course, the Discrete-Time Fourier Transform (DTFT)

$$G(\omega) = \sum_{-\infty}^{\infty} g[n] e^{-j\omega n}. \hspace{1cm} (2.48)$$

Figure 2.2: Example 2: two pulses (a), and (b), and their common autocorrelation function (c).
By replacing $e^{-jw}$ with $z$ we obtain

$$G(z) = \sum_{n=-\infty}^{\infty} g[n] z^{n},$$  \hspace{1cm} (2.49)

which is, of course, the usual $z$-transform of $g[n]$.* As before, $g[n]$ is assumed to have finite support. However, this time it is more convenient to assume that the support is bounded by 0 and $N$, therefore, the summing limits in the above formula should be replaced as follows

$$G(z) = \sum_{n=0}^{N} g[n] z^{n}.$$  \hspace{1cm} (2.50)

In this form the discrete $z$-transform looks almost exactly like the continuous Laplace transform (see Equation (2.8)). Similarly to Equation (2.19), it is easy to show that the $z$-transform of $g[n]$’s autocorrelation (denoted by $R(z)$) reads

$$R(z) = \overline{G(1/\bar{z})} G(z) = \bar{G}(z^{-1}) G(z).$$  \hspace{1cm} (2.51)

Note the different notation: $\overline{G(z)}$, and $\bar{G}(z)$, the former means that $G(z)$ is computed and then conjugated, the latter means that the coefficients of the polynomial $G(z)$ are conjugated. That is

$$\overline{G(z)} = \left( \sum_{n=0}^{N} g[n] z^{n} \right), \quad \bar{G}(z) = \sum_{n=0}^{N} \bar{g}[n] z^{n}.$$  \hspace{1cm} (2.52)

In other words, because $G(z)$ is a polynomial in $z$, it can always be presented as a product of simple factors

$$G(z) = A \prod_{k=1}^{N} (z - z_{k}),$$  \hspace{1cm} (2.53)

where $A$ is a scalar, and $z_{k}$’s are the roots of $G(z)$. Similarly,

$$\bar{G}(z^{-1}) = \bar{A} \prod_{k=1}^{N} (z^{-1} - \bar{z}_{k}).$$  \hspace{1cm} (2.54)

This yields

$$R(z) = |A|^{2} \prod_{k=1}^{N} (z - z_{k}) (z^{-1} - \bar{z}_{k}).$$  \hspace{1cm} (2.55)

*The notation where $G(z)$ is a polynomial in $z$ is common in geophysics. Electrical engineers, usually use $z^{-1}$ instead of $z$. 

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Obviously, both \( z_k \) and \( 1/\bar{z}_k \) are roots of \( R(z) \). Hence, if we consider a new signal \( g_2[n] \) whose \( z \)-transform is given by

\[
G_2(z) = G(z) \frac{z - 1/\bar{z}_k}{z - z_k} |z_k| ,
\]

where \( z_k \) is a non-zero root of \( G(z) \), then the auto-correlations of \( g_2[n] \) and \( g[n] \) will be equal, and so will be their power spectra. To show this, consider the \( z \)-transform of \( g_2[n]'s \) autocorrelation (denoted by \( R_2(z) \)):

\[
R_2(z) = \frac{G_2(1/\bar{z})G(z)}{(G(1/\bar{z}) - 1/\bar{z}_k) |z_k|} = R(z) \left( \frac{z - 1/\bar{z}_k}{z - z_k} \right) |z_k|^2 \]

By applying the inverse \( z \)-transform we immediately conclude that \( g_2[n] \) and \( g[n] \) have the same auto-correlation:

\[
r_2[n] \equiv g_2[n] \ast g_2[n] = g[n] \ast g[n] \equiv r[n] .
\]

This analysis leads to a simple conclusion: each non-zero and non-unitary (\( |z_k| \neq 1 \)) root of \( G(z) \) gives rise to two possible solutions whose auto-correlation is the same. Hence, for a general one-dimensional signal, whose support spreads over \( N+1 \) samples, there can be up to \( 2^N \) different signals* within the same support and same auto-correlation. Of course, similarly, to the continuous-time case that we considered in the previous section, in the case of real (and, probably, non-negative) signal \( g[n] \) the number of possible solutions may be smaller because not all roots of \( G(z) \) can be exchanged freely—some will result in complex (or, probably, negative) signals.

Another important observation is that if a solution \( x[n] \) has been bound, then all other solutions can be obtained from it by systematically replacing the roots of \( X[n] \), as described in Equation (2.55).*

Finally, recall that the sampling is done in the Fourier domain. Hence, to capture \( R(z) \) (the \( z \)-transform of \( g[n]'s \) auto-correlation), we must sample it at \( 2N+1 \) points. This requirement directly follows from the fact that \( R(z) \) is “almost a polynomial”, that is,

\[
R(z) = \frac{P_{2N}(z)}{z^N} ,
\]

*Here we do not count the trivial solutions that are obtained by multiplying by a constant phase factor.
where $P_{2N}(z)$ is a polynomial of degree $2N$. From Equation (2.55), we have

$$P_{2N} = |A|^2 \prod_{k=1}^{N} (z - z_k) (1 - z \bar{z}_k). \quad (2.60)$$

Due to the uniqueness of the interpolation polynomial (see, for example (Dahlquist & Björck, 2008)), it is sufficient to sample $P_{2N}$ at $2N + 1$ points to fully determine its coefficients, and, thus, to determine $R(z)$. For practical reasons the samples should be performed at the points that correspond to the DFT frequencies. Hereon we finish our treatment of the one-dimensional case and switch to multi-dimensional signals.

### 2.2 Multi-dimensional signals

The analysis in the two- or higher-dimensional case is very similar to what we have done in the one-dimensional case. The main result is a straightforward generalization of Equation (2.51). That is, given a two-dimensional time-discrete signal $g[n_1, n_2]$, whose support is given by $[0, N_1] \times [0, N_2]$, the $z$-transform of its autocorrelation is given by

$$R(z_1, z_2) = \overline{G(1/z_1, 1/z_2)G(z_1, z_2)} = \tilde{G}(z_1^{-1}, z_2^{-1})G(z_1, z_2), \quad (2.61)$$

where $G(z_1, z_2)$ is the two-dimensional $z$-transform of $g[n_1, n_2]$. From this formula we can easily find a way to generate a signal whose autocorrelation function is equal to that of $g[n_1, n_2]$. Let us assume that $G(z_1, z_2)$ can be represented as a product of two polynomials of lower degree:

$$G(z_1, z_2) = P(z_1, z_2)Q(z_1, z_2). \quad (2.62)$$

Assume further that the degree of $Q(z_1, z_2)$ in $z_1$ and $z_2$ is $d_1$ and $d_2$, respectively. Now we have

$$R(z_1, z_2) = \tilde{G}(z_1^{-1}, z_2^{-1})G(z_1, z_2)$$

$$= \tilde{P}(1/z_1, 1/z_2)\tilde{Q}(1/z_1, 1/z_2)P(z_1, z_2)Q(z_1, z_2)$$

$$= P(1/z_1, 1/z_2)Q(z_1, z_2)P(z_1, z_2)\tilde{Q}(1/z_1, 1/z_2)$$

$$= \tilde{G}_2(1/z_1, 1/z_2)G(z_1, z_2), \quad (2.63)$$

where

$$G_2(z_1, z_2) = P(z_1, z_2)\tilde{Q}(1/z_1, 1/z_2)z_1^{d_1}z_2^{d_2}. \quad (2.64)$$

Now, by applying the inverse $z$-transform to $G_2(z_1, z_2)$, we obtain a new signal $g_2[n_1, n_2]$ whose autocorrelation is equal to the autocorrelation of $g[n_1, n_2]$. Note that the multiplicative factor $z_1^{d_1}z_2^{d_2}$ makes $G_2(z_1, z_2)$ a proper polynomial in $z_1$ and $z_2$ whose degrees vary from 0 to $N_1$, and from 0 to $N_2$, respectively. Hence, it “shifts” $g_2[n_1, n_2]$ to the same support region $[0, N_1] \times [0, N_2]$ as occupied by $g[n_1, n_2]$. 

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Obviously $g_2[n_1, n_2] \neq g_1[n_1, n_2]$ whenever
\[ \bar{Q}(1/z_1, 1/z_2) z_1^{d_1} z_2^{d_2} \neq Q(z_1, z_2) . \] (2.65)

A similar result was obtained in (Hayes, 1982), however, the authors there considered only real-valued signals, and their approach was slightly different from ours.

So far, the development is essentially the same as we have seen in the one-dimensional case. The main result is also very similar: each factor of $G(z_1, z_2)$ can give rise to two different solutions (when inequality (2.65) hold). The main difference however, stems from the fact that multi-variate polynomials are, usually, irreducible, that is they cannot be factorized. More specifically, the set of reducible multi-variate polynomials is of measure zero. This fact was proved in (Hayes, 1982) for polynomials with real coefficients, however, its generalization to polynomials with complex coefficient is straightforward. In practical terms, this means that the chances of getting a reducible two- or three-dimensional polynomial are zero. This, in turn, means that the phase retrieval problem in the multi-dimensional case has, usually, a unique solution (up to the trivial transformations: lateral shifts, axis reversal, and constant phase factor).

Despite this “almost always” guaranteed uniqueness one must apply a critical judgment for every specific case, because physical signals may not be considered “purely random”. For example, the reducibility in the $z$-space has a clear physical meaning: if $G(z_1, z_2) = P(z_1, z_2) Q(z_1, z_2)$ then, $g[n_1, n_2] = p[n_1, n_2] \otimes q[n_1, n_2]$, where $\otimes$ denotes convolution. Hence, if the sought signal is a result of a convolution of some signal $p$ with a non-symmetric kernel $q$, the reconstruction will not be unique (even without counting the trivial transformations).
3 Current reconstruction methods

Current reconstruction methods date back to the pioneering work of Gerchberg and Saxton (GS) (Gerchberg & Saxton, 1972). Their original method was later improved significantly by Fienup (Fienup, 1982), who introduced the Hybrid Input-Output (HIO) algorithm. The latter algorithm, to the best of our knowledge, is the prevailing numerical method today for phase retrieval. The HIO method will be presented in Section 3.2. However, to better understand it, we start with its progenitor—GS, which is a classical example of optimization techniques known today as “alternating projections”. Details of the methods are give in Section 3.1 below. Before we proceed further, we need to define two basic terms:

**Definition 3.1.** The distance between a point \( x \) and a closed set \( S \) is defined as

\[
d(x, S) = \min_{y \in S} \|x - y\|. \tag{3.1}
\]

**Definition 3.2.** For a given point \( x \) and a closed set \( S \) we say \( y \in S \) is a projection of \( x \) onto \( S \) if:

\[
\|x - y\| = d(x, S). \tag{3.2}
\]

That is, \( y \) is a solution of the following minimization problem:

\[
\min_{y} \|x - y\|, \tag{3.3}
\]

subject to \( y \in S \).

It is important to note that a projection always exists, and furthermore it is unique if \( S \) is convex. Otherwise, there may be several solutions to Equation (3.3). As we shall see later, the constraints that appear in the phase retrieval problem are not convex. Nevertheless, the projection is still well defined (unique) in all cases, except when the current estimate has zeros in the Fourier domain. The non-convexity of the constraints along with existence of computationally cheap projections is the reason why current reconstruction methods are based on projections and why continuous optimization techniques, like gradient descent or Newton-type methods, are not capable of successful phase retrieval. More details on that will follow in Chapter 6. Meanwhile we proceed to the current reconstruction methods.
3.1 Gerchberg-Saxton method

Probably the first successful reconstruction method for phase retrieval was suggested by Gerchberg and Saxton for a slightly different problem—reconstruction of signals from two intensity measurements (Gerchberg & Saxton, 1972). The authors considered a situation that arises in electronic microscopes, where the intensity of the sought signal* can be measured along with its diffraction pattern (Fourier domain intensity). For this scenario, the authors suggested a reconstruction method that is based on projections (hereinafter the method will be referred to as GS method or GS algorithm). The algorithm is iterative, and each iteration consists of the following four steps:

**Step 1:** Fourier transform the current estimate of the signal.

**Step 2:** Replace the magnitude of the resulting computed Fourier transform with the measured Fourier magnitude to form a new estimate of the Fourier transform.

**Step 3:** Inverse Fourier transform the estimate of the Fourier transform.

**Step 4:** Replace the magnitude of the resulting computed signal with the measured signal modulus to form a new estimate of the signal.

It is a trivial exercise in basic calculus to show that the Steps 2 and 4 are, indeed, projections.

Of course, this algorithm can be (and, in fact, has been) generalized to a large number of situations, where the constraints in both domains are such that lead to a well defined and, preferably, computationally efficient projection. For example, this happens in the phase retrieval problem, where the object domain constraints are: limited support, that is, some parts of the signal are known to be zero; and, often, non-negativity—the signal in the support area is known to be real non-negative. A generalized version of GS is depicted in Figure 3.1 below.

*The signal is complex-valued, of course, otherwise the reconstruction would be trivial.*
It is important to stress that the imposition of the constraints (both in the Fourier and object domain) is performed via a projection. Projections, unlike general transformations, guarantee convergence of the algorithm as we prove below*. Before proceeding to the proof, note that convergence here means the lack of progress of the algorithm. It can happen in two different situations: first, arriving at some stationary point (a solution is always a stationary point, but not vice-versa); second, the algorithm can enter into an endless loop, jumping from point to point without decreasing the error.

**Theorem 3.1.** Let $S_O$ and $S_F$ be the sets of feasible signals as defined by the constraints in the object and Fourier domains, respectively. Furthermore, assume that these sets are equipped with the corresponding projection operators $P_O$ and $P_F$. Let $\{x^k\}_{k=0}^\infty$ and $\{y^k\}_{k=0}^\infty$ be the two sequences generated by the generalized GS method using the two projections:

$$y^k = P_F[x^k], \quad x^{k+1} = P_O[y^k]. \tag{3.4}$$

Then, the sequences $\{E_F^k\}_{k=0}^\infty$ and $\{E_O^k\}_{k=0}^\infty$, defined as

$$E_F^k = d(x^k, S_F), \quad E_O^k = d(y^k, S_O), \tag{3.5}$$

are monotonically decreasing, that is

$$E_F^{k+1} \leq E_F^k \tag{3.6}$$

$$E_O^{k+1} \leq E_O^k \tag{3.7}$$

*A similar theorem was proved by Fienup for a specific set of constraints (Fienup, 1982). Our proof is much more general.
Proof. Because $y^k$ is a projection of $x^k$ onto $S_F$ we have, by Definition 3.2, $d(x^k, S_F) = \|x^k - y^k\|$. Furthermore,

$$d(x^k, S_F) = \|x^k - y^k\| \geq \|x^{k+1} - y^k\|.$$  \hspace{1cm} (3.8)

Note that the inequality in the above equation follows from the fact that both $x^k$ and $x^{k+1}$ belong to $S_O$ and $x^{k+1}$ is a projection of $y^k$ onto $S_O$. Hence $\|y^k - x^{k+1}\| \leq \|y^k - x^k\|$. Similarly,

$$d(x^{k+1}, S_F) = \|x^{k+1} - y^{k+1}\| \leq \|x^{k+1} - y^k\|.$$  \hspace{1cm} (3.9)

Again, the inequality follows from the fact that both $y^k$ and $y^{k+1}$ belong to $S_F$ and $y^{k+1}$ is a projection of $x^{k+1}$ onto $S_F$. By combining Equations (3.8) and (3.9) we obtain

$$d(x^{k+1}, S_F) \leq \|x^{k+1} - y^k\| \leq d(x^k, S_F).$$  \hspace{1cm} (3.10)

Hence

$$d(x^{k+1}, S_F) \leq d(x^k, S_F).$$  \hspace{1cm} (3.11)

The proof for the second claim follows immediately if we write down Equation (3.10) for the iterations $k$ and $k + 1$:

$$d(x^{k+2}, S_F) \leq \|x^{k+2} - y^{k+1}\| \leq d(x^{k+1}, S_F) \leq \|x^{k+1} - y^k\| \leq d(x^k, S_F),$$  \hspace{1cm} (3.12)

and note that

$$\|y^k - x^{k+1}\| = d(y^k, S_O), \quad \|y^{k+1} - x^{k+2}\| = d(y^{k+1}, S_O).$$  \hspace{1cm} (3.13)

Thus, we obtain

$$d(y^{k+1}, S_O) \leq d(y^k, S_O).$$  \hspace{1cm} (3.14)

Note that all $x^k$ satisfy the object domain constraints and their discrepancy with the Fourier domain constraints is ever decreasing* with $k$. Similarly, all $y^k$ satisfy the Fourier domain constraints and their discrepancy with the object domain constraints is also ever decreasing. Hence, Theorem 3.1 may suggest that the GS method converges to a solution. This is true if the constraints are convex. In our case, however, the Fourier domain constraints are non-convex. Thus, the convergence to a solution is not guaranteed: the decrease in the functions $d(x^k, S_F)$ and $d(y^k, S_O)$ can be arbitrary small and even zero if the algorithm gets stuck at some stationary point (usually, a local minimum). Moreover, extensive experiments confirm that GS is not suitable for the standard phase retrieval from a single intensity measurement and support information (even for non-negative signals): the algorithm typically stagnates at some

*Strictly speaking, it is a non-increasing sequence, however, in practice most algorithms terminate after the decrease in the current step is below some threshold. Hence, the sequence is strictly decreasing during the algorithm execution.
point that is nowhere near a solution. In the next section we will review the Hybrid Input-Output algorithm that was invented by Fienup to overcome the stagnation problem of GS.

3.2 Fienup’s algorithms for phase retrieval

In 1982, Fienup suggested a family of iterative algorithms that are based on a different interpretation of the GS method (Fienup, 1982). These algorithms keep intact the right-hand side (Fourier domain) of the diagram depicted in Figure 3.1, that is, the first three operations of each iteration remains the same:

**Step 1:** Fourier transforming \( x^k \xrightarrow{\mathcal{F}} \hat{x}^k \).

**Step 2:** Satisfying the Fourier domain constraints \( \hat{x}^k \rightarrow \hat{y}^k \).

**Step 3:** Inverse Fourier transforming the result \( \hat{y}^k \xrightarrow{\mathcal{F}^{-1}} y^k \).

However, the further treatment (in the object domain) is different. Fienup’s insight was to group together the three steps above into a non-linear system having an input \( x \) and an output \( y \) as depicted in Figure 3.2. The useful property of this system is that the output \( y \) is always a signal having a Fourier transform that satisfies the Fourier domain constraints. Therefore, if the output also satisfies the object domain constraints, it is a solution of the problem. Unlike in the GS algorithm, the input \( x \) need no longer be thought of as the current estimate of the signal. Instead, it can be considered as a driving function for the next output \( y \). Hence, the input \( x \) need not necessarily satisfy the object domain constraints.

![Figure 3.2: Fienup’s interpretation of the Fourier domain constraints imposing procedure as a non-linear system.](image-url)
Based on this novel interpretation, Fienup suggested three algorithms for the phase retrieval problem from a single intensity and a priori knowledge that the signal $x$ is non-negative everywhere.

**input-output** This algorithm is based on a claim (see (Fienup, 1980)) that a small change of the input results in a change of the output that is a constant $\alpha$ times the change in the input. Hence, if a change $\Delta x$ is desired in the output, a logical choice for the change of the input to achieve that change in the output would be $\beta \Delta x$, where $\beta$ is a constant, ideally equal to $\alpha^{-1}$. For the problem of phase retrieval from a single intensity measurement the desired change of the output is

$$
\Delta x^k(t) = \begin{cases} 
0, & t \notin \nu, \\
-y^k(t), & t \in \nu,
\end{cases}
$$

(3.15)

where $\nu$ is the set of points at which $y(t)$ violates the object domain constraints. That is, at points where the constraints are satisfied, one does not require a change of the output. On the other hand; at points where the constraints are violated, the desired change of the output, required to satisfy the support and non-negativity constraints, is one that drives it towards the value of zero, and therefore, the desired change is the negated output at those points. Hence, the logical choice for the next input is

$$
x^{k+1}(t) = x^k(t) + \beta \Delta x^k(t)
= \begin{cases} 
x^k(t), & t \notin \nu, \\
x^k(t) - \beta y^k(t), & t \in \nu.
\end{cases}
$$

(3.16)

**output-output** This algorithm is based on the following observation with respect to the non-linear system depicted in Figure 3.2. If the output $y$ is used as an input, the resulting output will be $y$ itself, because it already satisfies the Fourier domain constraints. Therefore, irrespective of what input actually resulted in the output $y$, the output $y$ can be considered to have been resulted from itself as an input. From this point of view another logical choice for the next input is

$$
x^{k+1}(t) = y^k(t) + \beta \Delta x^k(t)
= \begin{cases} 
y^k(t), & t \notin \nu, \\
y^k(t) - \beta y^k(t), & t \in \nu.
\end{cases}
$$

(3.17)

Note that if $\beta = 1$ the output-output algorithm becomes the GS algorithm. Because best results are, in general, obtained with $\beta \neq 1$ the GS algorithm can be viewed as a sub-optimal version of the input-output algorithm.

*Originally, the algorithms were developed for real non-negative signals such as those that arise in astronomy. Later, the method was applied to complex-valued signals where it was discovered that precise support information is crucial for successful reconstruction (Fienup, 1987).
Finally, we consider the third algorithm suggested by Fienup. This time the next input is formed by a combination of the upper line of Equation (3.17) with the lower line of Equation (3.16):

\[ x^{k+1}(t) = y^k(t) + \beta \Delta x^k(t) = \begin{cases} 
    y^k(t), & t \notin \nu, \\
    x^k(t) - \beta y^k(t), & t \in \nu. 
\end{cases} \]

(3.18)

The last algorithm, known as the Hybrid Input-Output (HIO) algorithm, is currently the most widely used algorithm in industry due to its simplicity and (usually) best convergence rate amongst the above three algorithms.

In contrast to the GS method, there is no proof of convergence for the HIO method. However, a large body of experiments indicates that the algorithm is often successful in phase retrieval of real-valued non-negative signals. Still, stagnation is possible in certain situations (Fienup & Wackerman, 1986; Wackerman & Yagle, 1989). Besides, it turns out that phase retrieval of complex-valued objects whose support is not known precisely poses a much more severe problem. In this case, HIO is not capable of successful reconstruction (Fienup, 1987).

Several examples of reconstruction by HIO will be presented in the following sections. We do not present results obtained by the GS method because this method is not suitable for the phase retrieval problem—it stagnates very fast and its results are not even close to the sought signal.
4 Fundamental developments in optimization methods

Our research actually started with an idea to develop an efficient phase retrieval method based on continuous optimization technique. Unlike the current methods, described in the previous chapter, the continuous optimization approach can potentially provide significantly faster convergence rates and, probably even more important, can allow easy introduction of additional knowledge/assumptions into the computational scheme. The latter is especially difficult in the projection-based framework that is used in the current methods.

However, continuous optimization techniques, such as gradient descent or Newton-type methods, cannot be applied directly because the objective function \( f(z) \) is a real-valued function of complex variables: \( f: \mathbb{C}^n \to \mathbb{R} \). Therefore, its derivatives (of any order) with respect to \( z \) are not defined, as we show below. This can be circumvented by treating the real and the imaginary parts of \( z \) separately, that is, by looking at the function \( g: \mathbb{R}^n \times \mathbb{R}^n \to \mathbb{R} \), where \( z = x + jy \), and \( f(z) = f(z(x, y)) = g(x, y) \). This approach is viable and widely applied, though it may be more convenient to work with the original variable \( z \) (and its complex conjugate \( \bar{z} \)) rather than with its real and imaginary parts: \( x \), and \( y \). Moreover, because most modern computer languages provide native support for complex variables, this approach may be more efficient as well. Hence, in the subsequent sections we shall develop an alternative definition of the gradient and Hessian. Before that, let us demonstrate that \( f(z) \) is not differentiable with respect to \( z \), except in the trivial case where \( f(z) \) is constant.

**Lemma 4.1.** Let \( f(z) \) be a real function of complex argument \( z \), then \( f(z) \) cannot be holomorphic unless it is constant.

**Proof.** Let us denote the complex argument \( z = x + jy \) and \( f(z) = u(z) + jv(z) = u(x, y) + jv(x, y) \), where \( x, y, u, \) and \( v \) are real. If \( f(z) \) is holomorphic it must satisfy the Cauchy-Riemann equations

\[
\frac{\partial u}{\partial x} = \frac{\partial v}{\partial y}, \quad \frac{\partial u}{\partial y} = -\frac{\partial v}{\partial x}. \tag{4.1}
\]

However, \( f(z) \) is real, hence \( v(x, y) = 0 \) which, in turn means that

\[
\frac{\partial u}{\partial x} = \frac{\partial u}{\partial y} = 0. \tag{4.2}
\]

*The material presented in this section is currently in preparation for submission to a journal.*
Thus $u(x, y)$ is a constant and so is $f(z)$.

Before proceeding to the next section it is pertinent to define the following partial derivatives: $\frac{\partial f}{\partial z}$, and $\frac{\partial f}{\partial \bar{z}}$. Let $f(z) = h(z, \bar{z}) = u(x, y) + jv(x, y)$. If $f(z)$ is real, that is, if $v(x, y) = 0$, then

$$
\frac{\partial f}{\partial z} = \frac{1}{2} \left( \frac{\partial u}{\partial x} - j \frac{\partial u}{\partial y} \right),
\frac{\partial f}{\partial \bar{z}} = \frac{1}{2} \left( \frac{\partial u}{\partial x} + j \frac{\partial u}{\partial y} \right).
$$

The above derivatives are, sometimes, called the Wirtinger derivatives or Wirtinger operators (Wirtinger, 1927).

### 4.1 Complex gradient

Let us consider the first differential of a differentiable function $g : \mathbb{R}^n \mapsto \mathbb{R}$

$$
dg = \langle \nabla g, dx \rangle,
$$
where $\langle \cdot, \cdot \rangle$ denotes the usual inner product. This formula can be used for definition of a function’s gradient (see, for example, (Magnus & Neudecker, 1999)). However, this approach is not feasible in our case because the derivatives $\frac{\partial f}{\partial z_i}$ are not defined, unless $f(z)$ is holomorphic. And this, of course, is not possible except for some trivial cases where $f(z)$ is constant, as was shown in Lemma 4.1. Therefore, we suggest the following new definition for a real scalar function of a complex vector $f : \mathbb{C}^n \mapsto \mathbb{R}$

$$
df = \text{Re}\langle \nabla f, dz \rangle,
$$
where Re denotes the real part of a complex number. This definition preserves the most important properties of the gradient as we shall see later.

Now, to obtain an expression for $\nabla f$, we use an alternative form for the first differential using the partial derivatives $\frac{\partial f}{\partial z_i}, \frac{\partial f}{\partial \bar{z}_i}$, as was done in (Brandwood, 1983),

$$
df = (\nabla_z f)^T dz + (\nabla_{\bar{z}} f)^T d\bar{z},
$$
where

$$
\nabla_z f = \begin{bmatrix}
\frac{\partial f}{\partial z_1} \\
\vdots \\
\frac{\partial f}{\partial z_n}
\end{bmatrix}, \quad \nabla_{\bar{z}} f = \begin{bmatrix}
\frac{\partial f}{\partial \bar{z}_1} \\
\vdots \\
\frac{\partial f}{\partial \bar{z}_n}
\end{bmatrix}.
$$

That is, the function $f(z)$ is assumed to be a function of two independent vectors $z,$
\( \nabla \bar{z} f = \nabla \bar{z} f \).

Therefore, from Equation (4.6) we obtain

\[
\begin{align*}
    df &= (\nabla z f)^T dz + (\nabla \bar{z} f)^T d\bar{z} \\
    &= (\nabla z f)^T dz + (\nabla \bar{z} f)^T d\bar{z} \\
    &= 2 \text{Re} \left( (\nabla z f)^T dz \right) \\
    &= \text{Re} \langle 2\nabla z f, dz \rangle.
\end{align*}
\]

Hence, according to our definition in Equation (4.5), the gradient of \( f \) reads

\[ \nabla f(z) = 2\nabla \bar{z} f. \]  \( \text{(4.10)} \)

Note that Brandwood in his paper (1983) arrived at a slightly different definition: \( \nabla f(z) = \nabla \bar{z} f \), which is incorrect. However, being different by only the factor of two, it works in many situations because most algorithm use the gradient direction only to perform a line search, while its length is used exclusively as a termination criterion.

The following two theorems from (Brandwood, 1983) prove that both definitions are consistent with the main gradient properties used in optimization: (a) the gradient defines the direction of maximal ascent, and (b) the gradient being zero is a necessary and sufficient condition to determine a stationary point of \( f(z) \).

**Theorem 4.2.** Let \( f : \mathbb{C}^n \rightarrow \mathbb{R} \) be a real-valued scalar function of a complex vector \( z \). Let \( f(z) = h(z, \bar{z}) \), where \( h : \mathbb{C} \times \mathbb{C} \rightarrow \mathbb{R} \) is a real-valued scalar function of two complex vector variables and \( h \) is analytic with respect to \( z_i \) and \( \bar{z}_i \). Then either of the conditions \( \nabla z h = 0 \) or \( \nabla \bar{z} h = 0 \) is necessary and sufficient to determine a stationary point of \( f(z) \).

**Proof.** We can always express \( f \) as a function of \( 2n \) real variables \( x_k \) and \( y_k \), by using \( z_k = x_k + jy_k \): \( f(z) = u(x, y) \). Therefore, \( u(x, y) \) (and hence \( f(z) \)) is stationary if, and only if, \( \partial u/\partial x_k = \partial u/\partial y_k = 0 \) for all \( k \). From Equation (4.3) we immediately conclude that

\[
\begin{align*}
    \frac{\partial u}{\partial x_k} &= \frac{\partial u}{\partial y_k} = 0 \iff \frac{\partial h}{\partial z_k} \\
    \frac{\partial u}{\partial x_k} &= \frac{\partial u}{\partial y_k} = 0 \iff \frac{\partial h}{\partial \bar{z}_k} \quad (4.11)
\end{align*}
\]

Hence, \( f(z) \) has a stationary point if and only if \( \nabla z f = 0 \). Similarly \( \nabla \bar{z} f = 0 \) is also necessary and sufficient to determine a stationary point of \( f(z) \). \( \square \)
Theorem 4.3. Let $f(z)$ and $h(z, \bar{z})$ be two functions as defined in Theorem 4.2, then the gradient $\nabla f \equiv 2\nabla_{\bar{z}} f$ defines the direction of the maximal rate of change of $f$ with $z$.

Proof. Consider Equations (4.5) and (4.9) that define the gradient $\nabla f$. Obviously
\[ |df| = |\text{Re} \langle 2\nabla_{\bar{z}} f, dz \rangle| \leq |\langle 2\nabla_{\bar{z}} f, dz \rangle|. \tag{4.12} \]

Furthermore, according to the Cauchy-Schwarz inequality
\[ |\langle 2\nabla_{\bar{z}} f, dz \rangle| \leq \|2\nabla_{\bar{z}} f\| \|dz\|. \tag{4.13} \]

It is easy to verify that the equality in Equations (4.12) and (4.13) holds if, and only if, $\nabla_{\bar{z}} f = \alpha dz$ for some real positive scalar $\alpha$.

Note that the result of Theorem 4.3 follows from our definition of the gradient via the first differential: $df = \text{Re}(\nabla f, dz)$, and not from its particular formula.

4.2 Complex Hessian

The Hessian can also be obtained by treating the function $f : \mathbb{C}^n \mapsto \mathbb{R}$ as a function of $2n$ real variables that are the real and the imaginary part of $f$’s complex argument $f(z) = u(x, y)$. Then, using Equation (4.3), one can express it as partial derivatives with respect to $z$ and $\bar{z}$. However, this time the order of variables is more important.

For example, in (van den Bos, 1994), the author defines the following two vectors $v \in \mathbb{C}^{2n}$, $w \in \mathbb{R}^{2n}$
\[ v = \begin{bmatrix} z_1 \\ \bar{z}_1 \\ z_2 \\ \bar{z}_2 \\ \vdots \\ z_n \\ \bar{z}_n \end{bmatrix}, \quad w = \begin{bmatrix} x_1 \\ y_1 \\ x_2 \\ y_2 \\ \vdots \\ x_n \\ y_n \end{bmatrix}. \tag{4.14} \]

Using this definition, and the fact that
\[ \begin{bmatrix} z_n \\ \bar{z}_n \end{bmatrix} = \begin{pmatrix} 1 & j \\ 1 & -j \end{pmatrix} \begin{bmatrix} x_k \\ y_k \end{bmatrix}, \tag{4.15} \]
we immediately obtain
\[ v = Aw, \tag{4.16} \]
where $A$ is a block-diagonal matrix
\[ A = \text{diag} \left( \begin{pmatrix} 1 & j \\ 1 & -j \end{pmatrix} \right). \tag{4.17} \]
Hence van den Bos easily concludes that
\[ \nabla^2 w f = A^* (\nabla^2 \hat{z} f) A. \] (4.18)

Furthermore, by using \( A^{-1} = \frac{1}{2} A^* \), the relation can be reversed
\[ \nabla^2 v f = \frac{1}{4} A (\nabla^2 w f) A^*. \] (4.19)

However, we are not interested in the Hessian per se because we specifically aim for large-scale problems. Our goal is to find an expression for the Hessian-vector product. To this end we consider the first differential of the gradient (again, by treating \( z \) and \( \bar{z} \) as independent variables),
\[ d(\nabla f) = (\nabla^2 f) dz = (\nabla_z (\nabla f)) dz + (\nabla_{\bar{z}} (\nabla f)) d\bar{z}. \] (4.20)

Hence, multiplying a vector \( a \) with the Hessian \( \nabla^2 f \) reads
\[ (\nabla^2 f) a = (\nabla_z (\nabla f)) a + (\nabla_{\bar{z}} (\nabla f)) \bar{a}. \] (4.21)

In the next section we will see how to apply this formula to an objective function associated with phase retrieval.

### 4.3 Application to the phase retrieval problem

To exemplify our development with application to the phase retrieval problem, let us use the following objective function
\[ E(s) = \frac{1}{2} |||\hat{s}| - r|||^2, \] (4.22)
where \( \hat{s} \) denotes the Fourier transform of a signal \( s \), \( r \) denotes the measured magnitude of the Fourier transform, and \( || \cdot || \) denotes the standard \( l_2 \) vector norm. Note, that \( s \) and \( r \) are not necessarily one-dimensional vectors, hence, strictly speaking, the \( l_2 \) norm is not properly defined in all cases. A proper notation would be
\[ E(s) = \| \text{vec}(|\hat{s}| - r) \|^2, \] (4.23)
where the operator \( \text{vec}(\cdot) \) is a simple rearrangement of a multidimensional argument into a column vector in some predefined order. For example, let \( s \) be a two-dimensional \( m \times n \) signal (matrix) with \( s_i \) being its \( i \)-th column. Then, \( \text{vec}(s) \) is an \( mn \times 1 \) vector:
\[ \text{vec}(s) = \begin{bmatrix} s_1 \\ s_2 \\ \vdots \\ s_n \end{bmatrix}. \] (4.24)
Thus, in our convention the \text{vec()} operator transforms a matrix into a column vector by stacking the matrix columns. Of course, this operator is defined for signals of arbitrary (finite) dimensionality. For the sake of brevity, hereinafter we shall use \textit{s} and \textit{vec(s)} interchangeably and the appropriate form should be clear from the context. Let us now review the objective function defined by Equation (4.22)

\[ E(x) = \frac{1}{2} \| |\hat{s}| - r \|^2 \]

\[ = \frac{1}{2} \| |\mathcal{F}[s]| - r \|^2 \]

\[ = \frac{1}{2} \| |Fs| - r \|^2 . \]

(4.25)

Here, \( \mathcal{F}[s] \) denotes the Discrete Fourier Transform (DFT) operator applied to a (multidimensional) signal \textit{s}, and \( F \) represents the corresponding matrix, in the sense that

\[ \text{vec}(\mathcal{F}[s]) = F \text{vec}(s) . \]

(4.26)

We introduce the DFT matrix \( F \) just for mathematical notation. In practice, however, the DFT transform is performed by the Fast Fourier Transform (FFT) algorithm that never creates this matrix. Note also that \( Fs \) means, actually, \( F \text{vec}(s) \), however, the shorter notation is used, as we mentioned earlier. Consider now the final form of the objective function we obtained in Equation (4.25)—it can be viewed as a non-linear function of a complex argument \( z \equiv Fs \).

\[ E(s) = \frac{1}{2} \| |Fs| - r \|^2 = f(Fs) = f(z) . \]

(4.27)

Hence, \( f : \mathbb{C}^n \mapsto \mathbb{R} \), where \( n \) is the number of elements in \( Fs \) (equal to that in \textit{s}, of course). With the theory developed in the previous section we can now find the gradient of our objective function.

\[ d(E(s)) = df(z) \]

\[ = \text{Re} \langle \nabla f, dz \rangle \]

\[ = \text{Re} \langle \nabla f, d(Fs) \rangle . \]

(4.28)

\[ = \text{Re} \langle \nabla f, Fs \rangle \]

\[ = \text{Re} \langle F^* \nabla f, ds \rangle \]

Hence, using our definition we obtain

\[ \nabla E(s) = F^* \nabla f(z) , \]

(4.29)

where \( F^* \) denotes the Hermitian (conjugate) transpose of \( F \). Now, by using Equation (4.25), we have

\[ f(z) = \frac{1}{2} \| |z| - r \|^2 . \]

(4.30)
From which we obtain

\[ \nabla f = 2 \nabla \bar{z} f \]

\[ = 2 \nabla \bar{z} \left( \frac{1}{2} ||z| - r||^2 \right) \]

\[ = 2(|z| - r) \circ \nabla \bar{z} (|z|) \]

\[ = 2(|z| - r) \circ \nabla \bar{z} \left( (z \bar{z})^{\frac{1}{2}} \right) \]

\[ = 2(|z| - r) \circ z^{\frac{1}{2}} \circ \frac{1}{2} \bar{z}^{\frac{1}{2}} \]

\[ = \left( z - r \circ \frac{z^{\frac{1}{2}}}{\bar{z}^{\frac{1}{2}}} \right) \]

\[ = \left( z - r \circ \frac{z}{|z|} \right). \] (4.31)

Here \( \circ \) denotes the element-wise (Hadamard) product. Moreover, note that \( r \), and \( z \) are vectors, therefore quotients, and exponents, like \( z/|z| \), and \( z^{\frac{1}{2}} \), are assumed to be performed element-wise. This minor abuse of notation improves readability, therefore we use it instead of introducing some special notation. Substituting the above result into Equation (4.29) we obtain (by using \( z = Fs \))

\[ \nabla E(s) = F^* \nabla f(z) \]

\[ = F^* \left( z - r \circ \frac{z}{|z|} \right) \]

\[ = F^* \left( Fs - r \circ \frac{Fs}{|Fs|} \right) \] (4.32)

\[ = s - F^{-1} \left( r \circ \frac{Fs}{|Fs|} \right). \]

In this derivation we used the fact that \( F \) is unitary, therefore \( F^{-1} = F^* \). The expression for \( \nabla E(s) \) is remarkable because it bears a clear physical meaning, which will be discussed in the following chapters. Meanwhile we proceed with developments required for our optimization approach.

We already have the gradient of our objective function. Hence, we can deploy a variety of powerful optimization routines, such as Quasi-Newton methods. However, our choice should be limited to those that do not form a full approximation to the Hessian matrix because typical signals may easily contain \( 10^6 \)–\( 10^9 \) elements which renders the problem of Hessian storage too costly for a typical computer. In the following chapters we use the excellent Quasi-Newtonian method called L-BFGS, which uses limited memory to store an approximation to the Hessian matrix (Liu & Nocedal, 1989). However, to also allow for more powerful optimization methods we shall consider the second derivatives of the objective function. Our
main goal is to devise the Hessian-vector product formula that is used in many large scale optimization methods, for example, in the Conjugate Gradients (CG) method (Hestenes & Stiefel, 1952), and in the Sequential Subspace Optimization (SESOP) method (Narkiss & Zibulevsky, 2005). To this end we consider the first differential of the gradient $\nabla E(s)$

$$d(\nabla E) = d (F^* \nabla f)$$

$$= F^* d (\nabla f)$$

$$= F^* (\nabla^2 f) d z$$

$$= F^* (\nabla^2 f) d (Fs)$$

$$= F^* (\nabla^2 f) F ds.$$ 

(4.33)

Hence, according to the definition of the Hessian we get

$$\nabla^2 E(s) = F^* (\nabla^2 f(z)) F.$$ 

(4.34)

Recall that the Hessian $\nabla^2 f(z)$ has not been defined, instead we focus on the Hessian-vector product. Based on our development we can compute $(\nabla^2 E(s))a$ for any vector $a$

$$(\nabla^2 E(s))a = F^{-1} (\nabla^2 f(z)) Fa = F^* [((\nabla^2 f(z))(Fa)].$$ 

(4.35)

The brackets in the last expression are added to emphasize the order of efficient computation: first, the Fourier transform $\hat{a} = Fa$ is computed; second, the Hessian-vector product $(\nabla^2 f(z))\hat{a}$ is computed (described below); finally, the result undergoes an inverse Fourier transform. It is important to note that the first and the third steps in the above calculation are independent of the objective function and only the second step has this dependence. Let us now devise the formula for the Hessian-vector product $(\nabla^2 f(z))a$. Using Equation (4.21) we have

$$(\nabla^2 f) a = (\nabla_z (\nabla f)) a + (\nabla_{\bar{z}} (\nabla f)) \bar{a}$$

$$= \left( \nabla_z \left(z - r \circ z^\frac{1}{2} \circ \bar{z}^{-\frac{1}{2}} \right) \right) a + \left( \nabla_{\bar{z}} \left(z - r \circ z^\frac{1}{2} \circ \bar{z}^{-\frac{1}{2}} \right) \right) \bar{a}$$

$$= \text{diag} \left( 1 - \frac{r}{2|z|} \right) a + \text{diag} \left( r \circ z^2 \right) |2|z|^3 \bar{a}$$

(4.36)

$$= \left( 1 - \frac{r}{2|z|} \right) a + \left( r \circ z^2 \right) |2|z|^3 \bar{a}.$$ 

Note that we again use the quotient and exponent, like $r/z$ and $z^2$ in the element-wise manner.

### 4.3.1 Special properties

Let us consider some mathematical properties of the gradient and the Hessian of our objective function. First, let us look at the equation that defines the Newton
direction $d$

$$(\nabla^2 E) d = -\nabla E. \quad (4.37)$$

Even if we assume that the Hessian $\nabla^2 E$ is invertible, finding $d$ is not straightforward as we do not form $\nabla^2 E$ explicitly. Fortunately, the Hessian-vector product routine is sufficient. For example, we can use the CG method to find $d$. However, this will require a fair amount of iterations. To find a better (faster) way, let us consider the product $(\nabla^2 E) \nabla E$

$$(\nabla^2 E) \nabla E = F^* (\nabla^2 f(z)) F \nabla E$$

$$= F^* (\nabla^2 f(z)) FF^* \nabla f$$

$$= F^* (\nabla^2 f(z)) F \nabla f$$

$$= F^* (\nabla^2 f(z)) \left( z - r \circ \frac{z}{|z|} \right)$$

$$= F^* \left( \left( 1 - \frac{r}{2|z|} \right) \circ \left( z - r \circ \frac{z}{|z|} \right) + \frac{r \circ z^2}{2|z|^3} \circ \left( z - r \circ \frac{z}{|z|} \right) \right) \quad (4.38)$$

$$= F^* \left( z - \frac{r \circ z}{2|z|} - \frac{r \circ z}{|z|} + \frac{r \circ z}{2|z|^2} + \frac{r \circ z}{2|z|} - \frac{r \circ z}{2|z|^2} \right)$$

$$= F^* \left( z - \frac{r \circ z}{|z|} \right)$$

$$= \nabla E. \quad (4.38)$$

Namely, the gradient $\nabla E$ is an eigenvector of the Hessian $\nabla^2 E$ with the corresponding eigenvalue equal to one. This means that $-\nabla E$ is the Newton step. That is, the gradient descent method is equivalent to the Newton method in this case. Let us consider a single gradient descent (Newton) step with unit step-length

$$s - \nabla E(s) = F^{-1} \left( r \circ \frac{F_s}{|F_s|} \right). \quad (4.39)$$

Consider the above result from a physical point of view: the current signal estimate $s$ undergoes the Fourier transform $F_s$, then the (generally incorrect) magnitude $|F_s|$ is replaced with the correct one $r$, and the resulting signal is inverse transformed by $F^{-1}$. This is exactly the projection step that we saw in Chapter 3. Is then a single gradient descent step enough to solve the phase retrieval problem? The answer is yes, though the result is usually meaningless because it does not satisfy additional constraints that are usually imposed on the sought signal, for example, support information. The relation between the projection and the gradient descent has long been known (see (Fienup, 1982)), however, the relation to the Newton method is new, to the best of our knowledge.

We have found one eigenvalue (1) and eigenvector ($\nabla E$) of the Hessian. We may get even deeper insight into the problem if we look at the eigendecomposition of
the Hessian. To this end we need the full Hessian matrix. It can be obtained, using our Hessian-vector product, for the real-valued case, that is \( s \in \mathbb{R}^n \). Consider the Hessian-vector product for some real vector \( t \)

\[
(\nabla^2 E) t = F^* (\nabla^2 f) F t
\]

\[
= F^* (\nabla^2 f) (F t)
\]

\[
= F^* \left( \text{diag} \left( 1 - \frac{r}{2 |s|} \right) (F t) + \text{diag} \left( \frac{r \circ \hat{s}^2}{2 |s|^3} \right) (\overline{F} t) \right)
\]

\[
= F^* \left( \text{diag} \left( 1 - \frac{r}{2 |s|} \right) F + \text{diag} \left( \frac{r \circ \hat{s}^2}{2 |s|^3} \right) \overline{F} \right) t
\]

\[
= F^* \left( \text{diag} \left( 1 - \frac{r}{2 |s|} \right) F + \text{diag} \left( \frac{r \circ \hat{s}^2}{2 |s|^3} \right) F^* \right) t.
\]

Hence, we obtain

\[
\nabla^2 E = F^* \text{diag} \left( 1 - \frac{r}{2 |s|} \right) F + F^* \text{diag} \left( \frac{r \circ \hat{s}^2}{2 |s|^3} \right) F^*
\]

\[
= I - F^* \text{diag} \left( \frac{r}{2 |s|} \right) F + F^* \text{diag} \left( \frac{r \circ \hat{s}^2}{2 |s|^3} \right) F^*
\]

In this form, it is easy to perform an eigenanalysis of the Hessian. The main results are proven in the following two theorems.

**Theorem 4.4.** Let \( \nabla^2 E \) be as defined in Equation (4.41), where \( \hat{s} \) represents the Fourier transform of a real signal \( s \), and \( r \) denotes the absolute value of the Fourier transform of a real signal. Then, the eigenvalues of the Hessian are given by

\[
\lambda(\nabla^2 E) = 1 - \frac{r}{2 |s|} \pm \frac{r}{2 |s|}.
\]

**Proof.** To prove the claim we must recall certain properties of the DFT matrix \( F \).

1. \( F \) is unitary: \( F^* F = FF^* = I \).

2. \( F \) is symmetric: \( F^T = F \).

3. if \( t \) is real, then \( Ft \) is Hermitian, that is, conjugate symmetric.

4. \( F^2 \) is a permutation matrix that “reverses” its argument. As a consequence, if \( t \) is real, then \( F^2 (F t) = \overline{F} t \).

5. Applying the Fourier transform four times results in the original signal, namely \( F^4 = I \).
Now, let us consider a real signal $t$, and the following matrix $A$ obtained from it

$$
A = F^* \text{diag}(Ft) F^* \\
= F^* F^4 \text{diag}(Ft) F^4 F^* \\
= F^* F^2 F^2 \text{diag}(Ft) F^2 F^2 F^* \\
= F \text{diag}(Ft) F \\
= \overline{F} \text{diag}(Ft) \overline{F} \\
= \overline{F}^* \text{diag}(Ft) F^* \\
= \bar{A}.
$$

(4.43)

We find that $A$ is symmetric. Therefore, $A^2$ can be written as follows:

$$
A^2 = AA \\
= A \bar{A} \\
= F^* \text{diag}(Ft) F^* F \text{diag}(\overline{F}t) F \\
= F^* \text{diag}(|Ft|^2) F.
$$

(4.44)

Thus, the eigenvalues of $A^2$ are $|Ft|^2$, and the eigenvalues of $A$ are $\pm|Ft|$. Moreover, the matrix can be written as

$$
A = \sqrt{A^2} = F^* \text{diag}(\pm|Ft|) F.
$$

(4.45)

In this form, it is obvious that our proof is complete once we note that $(r \circ z^2)/(2|z|^3) = Ft$ for some real vector $t$. This is quite obvious, because $(r \circ z^2)/(2|z|^3)$ is Hermitian, hence the inverse Fourier transform will result in a real vector. \hfill \Box

Note that this proof does not tell us whether ‘+’ or ‘−’ should be taken in Equation (4.42), and it says nothing about the eigenvectors of the Hessian. These questions will be addressed in Theorem 4.6. However, this result can already provide some interesting insights into the problem. First, about half (see below) of the Hessian eigenvalues are unity (these correspond to the choice of ‘+’ in Equation (4.42)); the rest are equal $1 - r/|\hat{s}|$. Furthermore, if an exact solution is found, that is, if $|\hat{s}| = r$, then the eigenvalues of the Hessian become 1 and 0, with multiplicities equal, respectively, to the number of pluses and minuses in Equation (4.42). Hence, about half of the eigenvalues will be zero at a solution, which might make the problem quite difficult because the Hessian is highly singular at a solution and ill-conditioned in its neighborhood. Another observation shows that if $|\hat{s}| > r$ (the relation is taken element-wise), then the Hessian is positive definite, which is beneficial in optimization problem.

We next prove a theorem that is much stronger than Theorem 4.4. This time we devise an unambiguous expression for the Hessian eigenvalues and also find its eigenvectors. The complete derivation is split into the following two theorems.
Theorem 4.5. Let \( \hat{t} \in \mathbb{C}^n \) be the Fourier transform of a real-valued signal \( t \) (either one- or multi-dimensional): \( \hat{t} = Ft \). Let us also denote by \( K = \{k_1, k_2, \ldots, k_{n_1}\} \) and \( L = \{l_1, l_2, \ldots, l_{n_1}\} \) the two sets of indices that are exchanged upon an application of the permutation matrix \( F^2 \), while \( M = \{m_1, m_2, \ldots, m_{n_2}\} \), where \( n_2 = n - 2n_1 \), is the set of indices that are invariant under the permutation \( F^2 \). That is, if \( e_i \) is the \( i \)-th column of the identity matrix \( I_n \), then

\[
\begin{align*}
F^2 e_{k_i} &= e_{l_i}, & \forall i &\in 1, 2, \ldots, n_1, \\
F^2 e_{l_i} &= e_{k_i}, & \forall i &\in 1, 2, \ldots, n_1, \\
F^2 e_{m_i} &= e_{m_i}, & \forall i &\in 1, 2, \ldots n_2.
\end{align*}
\]  

(4.46)

Then, the eigenvalues of the matrix \( C = \text{diag}(\hat{t}F^2) \) are as follows

\[
\begin{cases}
\lambda_i = \hat{t}_i, & \text{if } i \in M, \\
\lambda_i = |\hat{t}_i|, & \text{if } i \in K, \\
\lambda_i = -|\hat{t}_i|, & \text{if } i \in L.
\end{cases}
\]  

(4.47)

The corresponding eigenvectors of \( C \) are given by

\[
\begin{align*}
v_{m_j} &= e_{m_j}, & \text{for } j &\in 1, 2, \ldots, n_2, \\
v_{k_j} &= e_{k_j} + a_{k_j}e_{l_j}, & \text{for } j &\in 1, 2, \ldots, n_1, \\
v_{l_j} &= e_{k_j} - a_{l_j}e_{l_j}, & \text{for } j &\in 1, 2, \ldots, n_1,
\end{align*}
\]  

(4.48)

where

\[
a_{k_j} = a_{l_j} = \frac{|\hat{t}_{k_j}|}{\hat{t}_{k_j}} = \frac{|\hat{t}_{l_j}|}{\hat{t}_{l_j}}.
\]  

(4.49)

Before we proceed to the proof, note that the set \( M \) is defined uniquely—it includes the zero-frequency and the half-Nyquist frequencies. The latter present if, and only if, the number of samples, along some dimension, is even. The two other sets: \( K \), and \( L \) are not unique—one can exchange \( k_j \) and \( l_j \). This non-uniqueness, however, does not have any special effect. The theorem, in fact, claims that the conjugate symmetric signal \( \hat{t} \) defines uniquely the set of eigenvalues and eigenvectors of \( C \) in the following manner. If \( \hat{t}_i \) has no conjugate counterpart (zero frequency, or half-Nyquist frequency) then it contributes the eigenvalue \( \hat{t}_i \) and the corresponding eigenvector \( e_i \).

If, on the other hand, \( \hat{t}_i \), \( i = k_j \) has a conjugate counterpart, then the pair \( \hat{t}_i = \hat{t}_{k_j} \), and \( \hat{t}_i = \hat{t}_{l_j} \) contributes two eigenvalues: \( |\hat{t}_i| \), and \( -|\hat{t}_i| \) along with the corresponding eigenvectors

**Proof.** Let us start with \( i \in M \):

\[
Ce_{m_j} = \text{diag}(\hat{t})F^2 e_{m_j}
= \text{diag}(\hat{t}) e_{m_j}
= \hat{t}_{m_j} e_{m_j},
\]  

(4.50)

40
which completes the proof for this case. Let us now consider \( i \in K \), that is, \( i = k_j \):

\[
\lambda_i v_i = \lambda_{k_j} v_{k_j} = Cv_{k_j} = \text{diag}(\hat{t}) F^2 (e_{k_j} + a_{k_j} e_{l_j}) = \text{diag}(\hat{t}) (F^2 e_{k_j} + a_{k_j} F^2 e_{l_j}) = \text{diag}(\hat{t}) (e_{l_j} + a_{k_j} e_{k_j}) = \hat{t}_{l_j} e_{l_j} + a_{k_j} \hat{t}_{k_j} e_{k_j} = \hat{t}_{k_j} e_{l_j} + a_{k_j} \hat{t}_{k_j} e_{k_j}.
\]

Hence, we have

\[
\lambda_{k_j} = a_{k_j} \hat{t}_{k_j},
\]

\[
\lambda_{k_j} a_{k_j} = \hat{t}_{k_j},
\]

which leads immediately to

\[
a^2_{k_j} \hat{t}_{k_j} = \hat{t}_{k_j} \Rightarrow a^2_{k_j} = \frac{\hat{t}_{k_j}}{t_{k_j}} \Rightarrow a_{k_j} = \pm \frac{|t_{k_j}|}{t_{k_j}}.
\]

To decide upon the sign of \( a_{k_j} \) in this equation, we use the fact that \( \lambda_{k_j} \) is positive, according to the definition in Equation (4.47). Now, if we look at Equation (4.52), we immediately obtain \( \lambda_{k_j} = a_{k_j} \hat{t}_{k_j} = \pm |t_{k_j}| \), hence, we must choose ‘+’. The proof for the last case, \( i \in L \), is very similar.

\[
\lambda_i v_i = \lambda_{l_j} v_{l_j} = Cv_{l_j} = \text{diag}(\hat{t}) F^2 (e_{l_j} - a_{l_j} e_{l_j}) = \text{diag}(\hat{t}) (F^2 e_{l_j} - a_{l_j} F^2 e_{l_j}) = \text{diag}(\hat{t}) (e_{l_j} - a_{l_j} e_{l_j}) = \hat{t}_{l_j} e_{l_j} - a_{l_j} \hat{t}_{l_j} e_{l_j} = \hat{t}_{l_j} e_{l_j} - a_{l_j} \hat{t}_{l_j} e_{k_j}.
\]

Hence, we have

\[
\lambda_{l_j} = -a_{l_j} \hat{t}_{l_j},
\]

\[
-\lambda_{l_j} a_{l_j} = \hat{t}_{l_j},
\]

which gives us

\[
a_{l_j} = \pm \frac{|\hat{t}_{l_j}|}{\hat{t}_{l_j}}.
\]
Again, we must choose ‘+’, because $\lambda_j = -a_j \check{t}_j = -|\check{t}_j|$ must be negative. It is worthwhile to note that the eigenvectors $v_i$ are mutually orthogonal.

Now we can prove the main result.

**Theorem 4.6.** Let $\hat{s}, \hat{t} \in \mathbb{C}^n$ be Fourier transforms of real-valued signals $s$, and $t$, respectively. Let the sets of indices $\mathcal{K}$, $\mathcal{L}$, and $\mathcal{M}$; and vectors $\{v_i\}_{i=1}^n$ be defined as in Theorem 4.5. Let $(\lambda_i, u_i)$ be an eigenpair of the following matrix

$$A = F^* \text{diag}(|\hat{s}|) F + F^* \text{diag}(\hat{t}) F^*.$$  \hspace{1cm} (4.57)

Then, the eigenvalues are given by

$$\begin{cases}
\lambda_i = |\hat{s}|_i + \hat{t}_i, & \text{if } i \in \mathcal{M}, \\
\lambda_i = |\hat{s}|_i - |\hat{t}|_i, & \text{if } i \in \mathcal{K}, \\
\lambda_i = |\hat{s}|_i + |\hat{t}|_i, & \text{if } i \in \mathcal{L}.
\end{cases} \hspace{1cm} (4.58)$$

And the eigenvectors are given by

$$U = [u_1, u_2, \ldots, u_n] = VF,$$ \hspace{1cm} (4.59)

where

$$V = [v_1, v_2, \ldots, v_n] \hspace{1cm} (4.60)$$

**Proof.** The proof is trivial once we note that

$$A = F^* \text{diag}(|\hat{s}|) F + F^* \text{diag}(\hat{t}) F^*$$

$$= F^* \text{diag}(|\hat{s}|) F + F^* \text{diag}(\hat{t}) F^4 F^*$$

$$= F^* \text{diag}(|\hat{s}|) F + F^* \text{diag}(\hat{t}) F^4 F^*$$

$$= F^* (\text{diag}(|\hat{s}|) + \text{diag}(\hat{t}) F^2) F$$

$$= F^* (B + C) F,$$

where

$$B = \text{diag}(|\hat{s}|), \quad C = \text{diag}(\hat{t}) F^2.$$ \hspace{1cm} (4.61)

The eigendecomposition of $C$ was found in Theorem 4.5. Thus, if we show that $v_i$ is also an eigenvector of $B$, with the corresponding eigenvalue of $|\hat{s}|_i$ the theorem will be proved. Indeed, it is easy to verify that

$$\text{diag}(|\hat{s}|) = |\hat{s}|_i v_i,$$ \hspace{1cm} (4.63)

which completes the proof. \hspace{1cm} $\Box$
Corollary 4.7. The eigenvalues of the Hessian $\nabla^2 E$ are given by

\[
\begin{cases}
\lambda_i = 1, & \text{if } i \in \mathcal{M}, \\
\lambda_i = 1 - \frac{r_i}{|x_i|}, & \text{if } i \in \mathcal{K}, \\
\lambda_i = 1, & \text{if } i \in \mathcal{L},
\end{cases}
\tag{4.64}
\]

where $\mathcal{M}$, $\mathcal{K}$, and $\mathcal{L}$ are set of indices, as defined in Theorem 4.5. The corresponding eigenvectors are given by $\{u_i\}_{i=1}^n$, as defined in Theorem 4.6.

This analysis can potentially lead to development of new methods. For example, because the eigenvalues of the Hessian is known, we can add a regularization term with a weight that guarantees that the problem is convex all the time. Although we have not developed such a method in the course of this work, it definitely appears on our “todo” list.

4.4 Concluding remarks (disappointing)

After developing the machinery for efficient optimization methods, we tested this apparatus on the classical phase retrieval problem for a real non-negative two-dimensional signal with known off-support locations ($\mathcal{O}$)

\[
\begin{aligned}
\min_x & \quad ||Fx - r||^2, \\
\text{subject to} & \quad x \geq 0, \\
& \quad x_{o\in\mathcal{O}} = 0.
\end{aligned}
\tag{4.65}
\]

Unfortunately, the classical Newton-type methods failed so solve this problem. This failure was not surprising because the phase retrieval problem is known to be “tough” for continuous optimization techniques. Let us cite a concluding excerpt from a study (Nieto-Vesperinas, 1986):

“The efficiency of an important class of Newton methods (the Levenberg-Marquardt algorithm) for solving overdetermined sets of nonlinear equations is tested in finding the solution to the two-dimensional phase problem. It is seen that the nonlinearity and number of local minima of the cost function increases dramatically with the size of the object array, making these methods of little practical use for sizes greater than $6 \times 6$. . . .”

Obviously, straightforward application of existing methods will not work. We hope that the eigenanalysis we performed in this chapter will eventually lead to a development of new efficient methods. Unfortunately, we did not do that in the framework of this research.


4.5 Concluding remarks (encouraging)

The failure of continuous optimization techniques when applied to the multi-dimensional phase retrieval problem is well known and, so to say, “widely accepted” amongst researchers. However, the exact reason for this is not fully understood. In Chapter 6 we provide an explanation for this failure. Moreover, in the following chapters we demonstrate that additional information can change things dramatically. For example, if a rough Fourier phase estimate is available, we demonstrate (first, empirically in Chapter 5, then theoretically in Chapter 6) that continuous optimization techniques succeed very well. Furthermore, in Chapter 6 we provide a rigorous mathematical reasoning that explains why any reasonable method is expected to succeed in the case where the Fourier phase uncertainty is below $\pi/2$ radians. In Chapter 9 we demonstrate that sparsity prior lets us solve successfully an even more difficult problem—simultaneous phase retrieval and bandwidth extrapolation.
5 Approximate Fourier phase knowledge for non-negative signals—first success∗

As was mentioned in Chapter 4, classical continuous optimization methods are known to fail miserably when applied to the phase retrieval problem. In Section 6 we shall give an explanation for this failure for a wide class of methods—monotone line-search optimization algorithms. In this chapter, however, we develop a new reconstruction method based on a Quasi-Newton optimization algorithm for a variation of the classical phase retrieval problem where very little additional information about the Fourier phase is available. In many situations, this information is readily available or can be obtained by an appropriate experimental arrangement.

A more extended discussion on some possible ways to obtain a rough phase estimate is delayed until Section 5.3. Here we demonstrate that a rough (up to $\pi$ radians) phase estimate allows us to develop a new method whose convergence rate is several orders of magnitude faster than that of the current reconstruction techniques. Unlike current methods, which are based on alternating projections, our approach is based on continuous optimization. Therefore, besides fast convergence, our method allows a great deal of flexibility in choosing appropriate objective functions as well as introducing additional information or prior assumptions about the sought signal like, for example, smoothness. The speed of convergence is important in many applications. For example, in microscopy a real-time algorithm would have a clear advantage. The ability to incorporate additional information may have an even greater effect: starting from a vast improvement in the reconstruction speed and going all the way to the very chances of successful reconstruction.

5.1 Developing an efficient optimization method

Let us start by formulating the optimization problem for real non-negative signals. The very common formulation is as follows

$$
\begin{align*}
\min_x & \quad \| |FPx|-r \|^2, \\
\text{subject to} & \quad x \geq 0,
\end{align*}
$$

(5.1)

∗The material presented in this chapter was published in (Osherovich et al., 2009b).
where $F$ denotes the Fourier transform operator (a matrix in the discrete case), $P$ represents zero-padding (note that any support information can be represented as zero-padding), and $r$ denotes the measured Fourier magnitude.

Of course, there is an endless number of ways to choose the objective function. The particular choice may affect the convergence speed and numerical stability. However, in our view, it is more important to choose the objective function that properly reflects the underlying physical phenomena. For example, the choice of Equation (5.1) is especially suitable when the measured quantity is $r$ and the noise in the measurements has a (close to) Gaussian distribution with zero mean.

As we have already seen, applying a Newton-type method to the problem in Equation (5.1) fails. However, we have not introduced yet the additional information available in the setup we consider here—the approximate Fourier phase. Let us consider one pixel in the Fourier domain. If the phase is known to lie within a certain interval $[\alpha, \beta]$, the correct complex number must belong to the arc $\hat{A}\hat{B}$ defined by $\alpha$ and $\beta$ as depicted in Figure 5.1a. Even with this additional information, the problem still remains non-convex and cannot directly be solved efficiently. However, if we perform a **convex relaxation**. That is, if we relax our requirements on the Fourier modulus and let the complex number lie in the convex region $\mathcal{C}$ defined by $\alpha$ and $\beta$ as shown in Figure 5.1b, the problem now becomes convex.

![Figure 5.1: Convex relaxation: (a) the original phase uncertainty interval results in an arc of a circle of known radius; (b) after relaxation, the allowed region is convex.](image)

Note that $\mathcal{C}$ is the smallest convex region that contains the original constraint (the arc $\hat{A}\hat{B}$). The formal definition of the relaxed problem is as follows:

\[
\begin{align*}
\min & \quad d^2(FPx, \mathcal{C}) \\
\text{subject to} & \quad x \geq 0
\end{align*}
\]

where $d(a, \mathcal{C})$ denotes the Euclidean distance from point $a$ to the convex set $\mathcal{C}$ (see
Definition 3.1). From our experience, a few dozen iterations are sufficient to solve this convex problem (see Figure 5.4a). Of course, the solution does not usually match the original image because both the phase and the magnitude may vary significantly. However, we suggest the following method for the solutions of the original problem.

**Stage 1:** Starting with a random $x^0$, solve the problem defined by Equation (5.2).

**Stage 2:** Use the solution obtained in Stage 1 (denoted $x^1$) as the starting point for the minimization problem that combines both the convex and non-convex parts, as defined below

$$
\begin{align*}
\min & \quad ||FPx - r||^2 + d^2(FPx, C) \\
\text{subject to} & \quad x \geq 0
\end{align*}
$$

(5.3)

More precisely, in our implementation we use the unconstrained minimization formulation, that is, instead of Equations (5.2) and (5.3) we minimize the following convex, and non-convex functionals, respectively.

$$
E_c(x) = d^2(FPx, C) + ||[x]_-||^2,
$$

(5.4)

$$
E(x) = ||FPx - r||^2 + \mu_1 d^2(FPx, C) + \mu_2 ||[x]_-||^2,
$$

(5.5)

where $[x]_-$ is defined as follows

$$
[x]_- = \begin{cases} 
0, & x \geq 0 \\
x, & x < 0
\end{cases}
$$

(5.6)

The weights $\mu_1$, and $\mu_2$ are usually set to unity. Results of our simulations are presented in the next section.

### 5.2 Simulations and Results

Due to the high dimensionality of the problem (especially in the 3D case) we limit our choice to methods that do not require the Hessian matrix or its approximation. Hence, in our implementation we use a modified version of the SESOP algorithm (Narkiss & Zibulevsky, 2005) and the L-BFGS method (Liu & Nocedal, 1989). Both algorithms demonstrate very similar results. The main difference is that SESOP guarantees that there are two Fourier transforms per iteration just like in the GS and HIO methods. The L-BFGS method, on the other hand, cannot guarantee that. However, in practice the average number of the Fourier transforms per iteration is very close to that of SESOP and HIO.

The method was tested across a variety of data. In this section we present some of these examples. The first example is a molecule of caffeine whose 3D model along with

*In fact, $x^0$ can be chosen in any reasonable way, as our method is insensitive to the choice of the starting point.*
a 2D projection of its electron density, and the corresponding Fourier magnitudes are shown in Figure 5.2. This information was obtained from a PDB* (protein database) file. In addition, we use a “natural” image which represents a class of images with rich texture and tight support. Moreover, it may be easier to estimate the visual reconstruction quality of such images. This image and its Fourier modulus are given in Figure 5.3.

Figure 5.2: Caffeine molecule: (a) 3D model (PDB), (b) 2D projection of its electron density; and their corresponding Fourier magnitudes: (c) 3D, and (d) 2D.

*See http://www.pdb.org for more information.
Figure 5.3: A natural image (Lena): (a) original image, and (b) its Fourier magnitude.

Note that we assume that a rectilinear sampling is available in the 3D case. In practice, however, the sensors measure a two-dimensional slice of the 3D volume. Provided that a sufficient number of such slices were measured, an interpolation can be used to form a rectilinear array of measurements (Miao et al., 2001). However, the slices can be incorporated directly into our minimization scheme. This will be addressed in future work.

In our experiments we tested a phase uncertainty of up to 3 radians. The bounds were chosen at random at every measured pixel (voxel) such that the true phase had a uniform distribution inside the interval. The starting point \( x^0 \) was also chosen randomly. Of course, there is an obvious way to make a more educated guess: by choosing the middle of the uncertainty interval, however, this choice will generally violate the object domain constraints. Fortunately, our experiments indicate that the starting point has little influence on the reconstruction. In all cases the reconstructed images obtained with our method were visually indistinguishable from the original. Therefore, we only present the values of \( E_c(x) \) and \( E(x) \) as defined in Equations (5.4) and (5.5) to visualize the progress of the first and the second stages, respectively. The second stage is compared with the HIO algorithm for which the error term is \( E(x) \) without the phase bounds constraint, that is,

\[
E_{\text{HIO}} = \|FPx - r\|^2 + \|[x]_-\|^2.
\]  

The first experiment is as follows. First, we run 60 iterations of Stage 1, that is, the convex problem defined by (5.2). The progress of different images is shown in Figure 5.4a. In the second stage we run 200 iterations of our algorithm (SESOP) starting at the solution obtained in the previous stage \( x^1 \). To compare the convergence rate with current methods, we ran twice the HIO algorithm: once, starting at \( x^0 \), whereby the algorithm is unaware of the additional phase information. Another run was started at \( x^1 \), hence, the phase information was made (indirectly) available to the algorithm. The results for 2D and 3D reconstruction of the caffeine molecule
are shown in Figures 5.4b and 5.4c, respectively. The results of the natural image are shown in 5.4d.

It is evident from these results that our method significantly outperforms the HIO algorithm in all experiments. Moreover, its superiority for the “Lena” image is tremendous.

![Figure 5.4: Reconstruction results: (a) stage 1 convergence rate, (b) stage 2 convergence rate of the 2D caffeine model, stage 2 of the 3D caffeine model, and (d) stage 2 of Lena.](image)

In addition to the examples shown in this chapter we have studied a number of other examples. Based on our observations we conclude that our algorithm demonstrates a significantly better convergence rate so long as the interval of phase uncertainty is not too close to $\pi$ radians.

Besides the fast convergence rate, our method allows us to incorporate additional information about the image or the noise distribution in the measurements. For example, in practice we measure $r^2$ and not $r$, and the noise distribution is Poissonian rather than Gaussian. In this case the maximum-likelihood criterion implies the functional for the error measure in the Fourier domain to be as follows:

$$E_P(x) = 1^T \left( |FPx|^2 - r^2 \ln (|FPx|^2) \right).$$  \hspace{1cm} (5.8)

To demonstrate the performance of our method we contaminated the measurements
Figure 5.5: Reconstruction from noisy data: (a) our method (30dB), and (b) HIO (16.7dB).

\(r^2\) of the “Lena” image with Poissonian noise such that the signal to noise ratio (SNR) was 53.6 dB. The phase uncertainty was 3 radians as before. First, we started by solving the convex problem, as defined by Equation (5.2). The solution obtained was then used as the starting point for the second stage of our method using the non-convex functional defined in Equation (5.8). The HIO algorithm also started at this solution. In addition to using the objective function that fits the noise distribution we also included a regularization term in the object space. In this example, we used the total variation functional (Rudin et al., 1992)

\[
TV(x) = \int |\nabla x|,
\]

with a small weight. Total variation is a good prior for a broad range of images, especially for images that are approximately piece-wise constant. In our case, introduction of this regularization added about 3 dB to the reconstruction SNR. The reconstruction results are shown in Figure 5.5. Our method achieved the SNR of 30dB, while the HIO algorithm produced a significantly inferior result. Its SNR was only 16.7dB. Note that the SNR values given above were obtained using different measures. The SNR in the measurements was obtained with respect to the measured intensity, that is, squared Fourier magnitude, while the SNR values reported for the reconstruction results was measured with respect to the image magnitude. More correct would be to measure it with respect to the image intensity—in this case the reported SNR should be multiplied by approximately 2. Hence, our reconstruction provides SNR that is better than the SNR of the measurement. This improvement was achieved with the aid of the regularization term (TV)—something that is impossible to incorporate into methods based on alternating projections.

It is also worthwhile to note that Poissonian noise of small intensity can be well approximated by Gaussian noise. However, if one uses the objective function implied
by the Gaussian noise in $r^2$, that is,

$$\| |FPx|^2 - r^2\|^2,$$  \hspace{1cm} (5.10)

the reconstruction results are a few dB’s worse than those we got with the proper choice of the objective function.

### 5.3 Concluding remarks

In this chapter we presented the first successful method based on continuous optimization for the phase retrieval problem for non-negative objects whose phase is known to lie within a certain interval. It is important to note, however, that straightforward incorporation of this information does not lead automatically to a successful method of reconstruction. Therefore, we designed a two-stage algorithm. At the first stage we perform convex relaxation and solve the resulting convex problem. At the second stage the original objective function is re-introduced into the scheme and the reconstruction continues from the solution of the first stage. The algorithm demonstrates a significantly better convergence rate compared to current reconstruction methods. Moreover, in contrast to these methods, our technique is flexible enough to allow incorporation of additional information. Practical examples of such information include measurement noise distribution and knowledge that the sought image is piece-wise smooth.

It is worthwhile to discuss possible sources of the approximate Fourier phase information. Probably, most obvious way to obtain it is to introduce into the scene an object whose Fourier transform is known. In this case the recorded data is the modulus of a sum of two complex numbers, one of which is known, and we are actually required to perform some sort of holographic/interferometric reconstruction. However, unlike in the classical holography interferometry, the known object must not be known precisely. This is a very important property, because calibrating and maintaining an interferometer so as to keep the reference beam with high precision during a series of experiments is a time-consuming and expensive procedure. We shall return to the holographic setup in Chapters 7 and 8. But there are other sources of the phase information that do not require a physical modification of the experiments. For example, as explained in (Goodman, 2005, Section 5.2), an ideal lens can perform the Fourier transform. This property can be used to test lenses by illuminating them with a known beam and measuring the resulting intensity in the Fourier plane. By reconstructing the illuminating beam from this intensity and comparing it with the actual beam, one can estimate the quality of the lens. Note that the imperfections in the lens production affect directly the Fourier phase by diverting it from the known expected values. Another possible way to obtain a phase estimate is to use current reconstruction methods such as HIO in situations where they are known to be able to reconstruct the sought signal. If a successful reconstruction is guaranteed, at some point the phase must be close enough to the true value, and if we manage to identify

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such a moment, then we can switch to our algorithm and get a faster reconstruction. As a last resort, the whole interval of $2\pi$ radians can be split into several, small enough, intervals and each one can be tested separately. In this case, once a correct partitioning is found our method will recover the image. This approach is closely related to combinatorial optimization, as we have to choose a correct combination of the phase intervals. Thus it is very important that an assessment of the current hypothesis (a particular choice of the phase intervals) can be performed very fast.
6 Approximate Fourier phase data for arbitrary signals—why does it help?*

In the previous chapter we presented an algorithm for fast phase retrieval in situations where a rough Fourier phase estimate is available. Our experiments demonstrated excellent convergence rates for phase uncertainty intervals of up to \( \pi \) radians. This algorithm is potentially important, as it is the first successful application of continuous optimization technique to the phase retrieval problem. However, these results were purely empirical—no explanation has been provided so far about why and when this method is expected to solve the problem. Moreover, all these results were obtained for real-valued non-negative signals. When we tried to apply the method to complex-valued signals the results were less encouraging. The reason for this gap lies in our experimental setup: the phase uncertainty intervals were generated independently for each pixel in the Fourier space, ignoring the conjugate symmetry exhibited by real-valued signals in the Fourier domain. Hence, the effective phase uncertainty in our previous experiments was approximately \( \pi/2 \) radians and not \( \pi \). After we refined the allowed phase uncertainty interval, the results in both the real-valued and complex-valued cases returned to be excellent. This chapter provides an explanation for this phenomenon.

Thus, in this section we continue to evaluate the importance of approximate phase information in the phase retrieval problem. However, now we address more theoretical questions, the main one being when and why a rough phase estimate (up to \( \pi/2 \) radians) can lead to a guaranteed reconstruction by the algorithm presented in the previous section. Our main discovery is that the above phase uncertainty limit practically guarantees a successful reconstruction by any reasonable algorithm for the reason described below. This is an important property as it allows development of very efficient algorithms whose reconstruction time is orders of magnitude faster than that of the current method of choice—the Hybrid Input-Output (HIO) algorithm. We have already presented one such algorithm in the previous section, however, we believe that its results can be further improved by using more sophisticated interior point methods. Using the new algorithms we were able to reconstruct signals that cannot be successfully reconstructed by HIO, namely, complex-valued signals without tight support information.

Additionally, we provide a heuristic explanation of why continuous optimization

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*The material presented in this chapter was published in (Osherovich et al., 2011).
methods such as gradient descent or Newton-type algorithms fail when applied to the phase retrieval problem and how the approximate phase information can remedy this situation. We actually show the reason for the failure for a very large family of optimization methods—monotone line-search optimization algorithms. Notwithstanding this failure in the general case, we argue that a rough phase estimate leads to an important property: local minima of a functional associated with the phase retrieval problem are likely to be global minima. This is the reason for our previous claim: chances are that any algorithm capable of finding a local minimum will successfully reconstruct the image.

Additional numerical simulations are provided in Section 6.2 to demonstrate the validity of our analysis and success of our reconstruction method.

6.1 Optimization methods: the problem and the remedy

Let $z$ represent the true (unknown) signal, with $\hat{z}$ being its Fourier transform, in accordance with our notational conventions. The current estimate (obtained after $k$ iterations) is denoted by $x^k$. With this notation, the classical phase retrieval problem reads: find $x$ such that $|Fx| = |\hat{z}|$ subject to certain constraints in the object domain. When the Fourier phase is known to lie within a given interval $[\alpha, \beta]$ ($\alpha$, and $\beta$ are vectors of appropriate size) the problem to be solved is: find $x$ such that $|Fx| = |\hat{z}|$ and $\alpha \leq \angle(Fx) \leq \beta$ subject to certain constraints in the object domain. Note that such $x$ is not necessarily equal to $z$; even when the Fourier domain data is sufficiently “oversampled”, the reconstruction in the classical phase retrieval problem is not unique (Hayes, 1982), and it may not be unique even when the Fourier phase estimates are available.

Let us now try to understand why the classical Newton-type and gradient descent methods fail for the phase retrieval problem. Actually, we can address a very wide family of optimization methods that includes these two methods—monotone line-search algorithms. Each iteration of these algorithms has a common 3-step template:

**Step 1:** Find a descent direction $p^k$.

**Step 2:** Along that direction find a step-length $\Delta^k$ that sufficiently decreases the objective function value.

**Step 3:** Move to the new location: $x^{k+1} = x^k + \Delta^k p^k$.

Descent direction is defined as a vector whose inner product with the gradient is negative (for obvious reasons). This guarantees that there always exists a step along that direction that decreases the objective function value.

To be specific we choose the most popular objective function for the discrepancy minimization in the Fourier domain:

$$f(x) = \frac{1}{2} \| |Fx| - |\hat{z}| \|^2 .$$

(6.1)
The gradient of \( f(x) \) is given by (see Chapter 4):
\[
\nabla f(x) = x - F^{-1}\left( |\hat{z}| \circ \frac{Fx}{|Fx|} \right).
\]
(6.2)

As before, \( a \circ b \) and \( \frac{a}{b} \) denote the Hadamard (element-wise) product and quotient, respectively. It is interesting to note that the signal \( F^{-1}\left( |\hat{z}| \circ \frac{Fx}{|Fx|} \right) \) has a clear physical meaning: it is obtained from \( x \) by substituting the (wrong) Fourier magnitude \( |Fx| \) with the correct one \( |\hat{z}| \). Thus, it is nothing but the signal denoted by \( y \) in Figure 3.2. However, the main observation about the gradient is that \( \nabla f(x) = 0 \) if and only if \( |Fx| = |\hat{z}| \), that is, if and only if \( x \) is a solution. Of course, this is true only if there are no additional constraints. In practice, however, the optimization is done while respecting certain restrictions on \( x \). The constraints are often implemented as penalty functions that augment the original functional, and the augmented gradient may vanish when \( |Fx| \neq |\hat{z}| \). Also, we may find ourselves in a situation where no feasible descent direction exists, if the constraints are kept as ideal barriers. Such situations usually indicate a local minimum and make further progress by such standard optimization methods impossible. In this discussion we are being deliberately vague about the exact nature of the object domain constraints and enforcement thereof in the optimization process. We only assume that imposing these constraints on an image estimate will take that estimate to be closer\(^*\) to the true signal—a natural assumption for monotone optimization.

Let us now consider a single element in the Fourier domain. Using our notation, the true value is \( \hat{z}_i \), whose magnitude \( |\hat{z}_i| \) is known and whose phase \( \theta_i \) is unknown. We distinguish three possible scenarios where \( |\hat{x}_i| \neq |\hat{z}_i| \). First, the Fourier magnitude of the current estimate is smaller than \( |\hat{z}_i| \) and the phase error \( \alpha_i \) is greater than \( \pi/2 \) radians. Second, the current Fourier magnitude is greater than \( |\hat{z}_i| \) (phase error is unimportant in this case). Finally, the third possibility: the current estimated Fourier magnitude is less than \( |\hat{z}_i| \) and the phase error \( \alpha_i \) is less than \( \pi/2 \) radians. These scenarios are illustrated in Figures 6.1a, 6.1b, and 6.1c, respectively.

\(^*\)“Closer” here refers to the Euclidean distance, however it can be readily generalized to other metrics.
Recall also that, by Equation (6.2), we know that the gradient descent direction is always taking us from $\hat{x}$ towards $\hat{y}$ (see Equation (6.2) and Figure 6.1). Let us consider the first case. In the Fourier domain, when we move from $\hat{x}_i$ towards $\hat{y}_i$ we are actually moving away from the correct value $\hat{z}_i$ (see Figure 6.1a). Therefore, due to the unitarity of $F$, $x$ is pulled farther away from $z$. On the other hand, object domain constraints shall pull us towards the correct value, as assumed before. Hence, the two forces may cancel each other, resulting thereby in the stagnation of the algorithm. In numerical tests this stagnation is observed in all but very tiny problems. Worse still, it happens at very early stages, long before the reconstruction algorithm
gets close to the correct value. Hence, the results are usually worthless. In the second case, where the magnitude of the current estimate is greater than the correct value $\hat{z}_i$, moving from $\hat{x}_i$ towards $\hat{y}_i$ will necessarily bring us closer to the correct value $\hat{z}_i$ (see Figure 6.1b). The last case is more interesting. If the estimated Fourier magnitude is sufficiently smaller than the correct one, and the phase error $\alpha_i$ is less than $\pi/2$ radians, then moving along the gradient descent direction, that is, from $\hat{x}_i$ towards $\hat{y}_i$, will get us closer to $\hat{z}_i$, so long as we do not pass the point $\hat{z}_i'$ which is the projection of $\hat{z}_i$ onto the line segment $[0, \hat{y}_i]$ as shown in Figure 6.1c. Note that the fact that moving from $\hat{z}_i'$ towards $\hat{y}_i$ takes us farther away from the correct value $\hat{z}_i$ keeps us from claiming that any optimization algorithm will necessarily converge to a solution. However, in the discussion that follows we prove that the situation where an optimization algorithm gets stuck at some $x$ such that all $\hat{x}_i$ lie between $\hat{z}_i'$ and $\hat{y}_i$ is impossible. In fact, we argue below that in the situation where the Fourier phase errors are limited by $\pi/2$ radians any local minimum is likely to be a global one. That is, any algorithm capable of finding a (constrained) local minimum can be expected to solve the phase retrieval problem in this case. To give a reasoning behind this claim we must be more specific about the constraints in the object domain. As will be evident from the argument below, our assumptions are very general and fit all commonly encountered cases.

Let us consider the most frequent object domain constraint: limited support information.

$$x_o = 0, \quad \forall o \in O,$$

(6.3)

where $O$ denotes the set of off-support locations, where $z$ is known to be zero. Note that zero-padding is a special case of such support information. In certain situations the sought signal can be assumed to be real non-negative. In these situations the above constraint can be extended by the non-negativity requirement:

$$x_s \geq 0, \quad \forall s \not\in O.$$

(6.4)

What is important for our discussion is that in both cases the set of all feasible signals ($Z$) is a convex set that contains a proper cone $K$. That is, if $x$ and $y$ are feasible, then $\lambda x + \gamma y$ is also feasible for any non-negative scalars $\lambda, \gamma$. $Z$ contains $K$ but not necessarily equals it, but this is unimportant in the following discussion. Let us now consider the minimization of the objective function from Equation (6.1) subject to the following two conditions: first, the set of object domain constraints is convex and contains the proper cone $K$; second, the phase error (of all elements) in the Fourier domain is bounded by $\pi/2$, that is, $|\angle(\hat{x}) - \angle(\hat{z})| \leq \pi/2 - \epsilon$ for some small positive $\epsilon$. Obviously, the Fourier domain constraints define a convex set which is a proper cone too (a Cartesian product of proper cones). Hence, the optimization in this case is done over a convex set. Assume now that the algorithm converges at some local minimum $x$ which is not a solution (global minimum). Following a basic theorem from constrained-optimization theory we conclude that the following inequality must hold:

$$x_s \geq 0, \quad \forall s \not\in O.$$
hold for any feasible point \( w \) (see, for example, (Bertsekas, 1999)):

\[
\langle \nabla f(x), (w - x) \rangle \geq 0.
\] (6.5)

In words, that means that no feasible descent direction exists at the point \( x \). Let us consider a (small) subset of all feasible points: \( w = \lambda z + \gamma x \), where \( \lambda, \gamma \geq 0 \) (note that \( x \) and \( z \) are feasible by definition, hence, \( x, z \in K \) which implies that \( w \in K \)). The choice of this subset of feasible directions is stipulated by the fact that the phase error of \( w \) in all frequencies is less than or equal to the phase error of \( x \) (it is strictly smaller when \( \lambda \neq 0 \)). With \( w \) as above, Equation (6.5) becomes

\[
\langle \nabla f(x), (w - x) \rangle = \lambda \langle \nabla f(x), z \rangle + (\gamma - 1) \langle \nabla f(x), x \rangle.
\] (6.6)

If \( \langle \nabla f(x), x \rangle < 0 \), we can set \( \lambda = 0 \), \( \gamma = 2 \) and get a feasible descent direction (this is, actually, scaling up of \( x \)). If \( \langle \nabla f(x), x \rangle > 0 \) we can set \( \lambda = 0 \), \( \gamma = 0.5 \) and, again, get a feasible descent direction (scaling down of \( x \)). Hence, any local minimum must obey \( \langle \nabla f(x), x \rangle = 0 \) (we shall call such \( x \) optimally scaled). In this situation, the sign of the left-hand size of Equation (6.5) depends solely on the sign of the inner product \( \langle \nabla f(x), z \rangle \). It is more convenient to consider the above inner product in the Fourier domain. Due to unitarity of \( F \) we have:

\[
\langle \nabla f(x), z \rangle = \langle F \nabla f(x), Fz \rangle
= \sum_i (|\hat{x}_i| - |\hat{z}_i|) |\hat{z}_i| \cos \alpha_i.
\] (6.7)

Recall that the above formula is considered in the context of an optimally scaled \( x \), that is:

\[
0 = \langle \nabla f(x), x \rangle = \langle F \nabla f(x), Fx \rangle
= \sum_i (|\hat{x}_i| - |\hat{z}_i|) |\hat{x}_i|.
\] (6.8)

For our following discussion, it is convenient to consider Equations (6.8) and (6.10) as weighted sums of \( |\hat{x}_i| - |\hat{z}_i| \). Thus, for example, it becomes obvious that an \( x \) that belongs to the convex region \( C \) (see Figure 5.1b), as was required in the algorithm in Chapter 5, cannot be a local minimum unless \( |\hat{x}| = |\hat{z}| \) (which makes it a global one), because this is the only way to get zero by summing non-positive numbers associated with strictly positive weights. This explains the success of the algorithm. However, even without the restrictions on \( |\hat{x}| \) used in the original algorithm, we can expect the sum in Equation (6.8) to be negative. To understand why let us split it into three
disjoint sets of indices,

\[ \sum_i (|\hat{x}_i| - |\hat{z}_i|) |\hat{z}_i| \cos \alpha_i = \sum_{i \in B} (|\hat{x}_i| - |\hat{z}_i|) |\hat{z}_i| \cos \alpha_i + \sum_{i \in S} (|\hat{x}_i| - |\hat{z}_i|) |\hat{z}_i| \cos \alpha_i + \sum_{i \in A} (|\hat{x}_i| - |\hat{z}_i|) |\hat{z}_i| \cos \alpha_i, \quad (6.11) \]

where \( B = \{ i \mid |\hat{x}_i| > |\hat{z}_i|\} \), \( S = \{ i \mid |\hat{x}_i| \leq |\hat{z}_i| \cos \alpha_i \} \), and \( A = \{ i \mid |\hat{z}_i| \cos \alpha_i < |\hat{x}_i| < |\hat{z}_i|\} \). With this subdivision it is easy to compare the sum in Equation (6.8) (for which a negative sign indicates the presence of a feasible descent direction) and the sum in Equation (6.10) (which is zero). The weight in these weighted sums changes from \(|\hat{x}_i|\) in (6.10) to \(|\hat{z}_i| \cos \alpha_i\) in (6.8). Hence, for \( i \in B \), \(|\hat{x}_i| - |\hat{z}_i|\) is positive and its weight has decreased, thus, pulling the total sum towards a negative value. For \( i \in S \), \(|\hat{x}_i| - |\hat{z}_i|\) is negative, and its weight has increased, thus again contributing to the negativeness of the result. The only subset of indices that increases the sum is \( i \in A \). From our experience it is very unlikely to encounter a situation where the contribution of \( i \in A \) outweighs the joint contribution of \( i \in B \) and \( i \in S \). Hence it is very unlikely to get stuck in a local minimum with no descent direction. Note also that if the phase error of all frequencies of \( x \) is strictly less than \( \pi/2 \) radians then the sum in Equation (6.8) must be zero for \( x \) to be a local minimum, because if the direction towards \( w \) is an ascent direction, one can simply reverse it to get a descent direction.

This discussion provides a heuristic explanation why a carefully designed optimization method can be expected to converge to a global minimum, that is, to solve the phase retrieval problem when the Fourier phase is known up to \( \pi/2 \) radians and the object domain constraints are given in the form of (possibly loose) support information.

### 6.2 Experimental results

The method was tested on many images with consistent results. Here, for demonstration purposes, we chose a natural image with a lot of features so as to allow easy perception of the reconstruction quality by the naked eye. We demonstrate our results on two different cases: one with loose support information, that is, part of the image is zero and we do not know that a priori; another with tight support. Both images are complex-valued and their original size is \( 128 \times 128 \) pixels, padded with zeros to the size of \( 256 \times 256 \) pixels. The intensity (squared magnitude) of the images (without the zero-padding) is shown in Figure 6.2.
All our experiments show that the phase distribution in the object domain does not affect the reconstruction. Therefore, the actual phase distribution was chosen to be random to avoid any possible assumptions of smoothness. We compared three reconstruction methods. First, a slight modification of the quasi-Newton method from Chapter 5. Second, we created the following phase-aware modification of the HIO algorithm (PA-HIO). When the Fourier phase is known to lie in the interval $[\alpha, \beta]$, PA-HIO’s correction in the Fourier domain forces the current estimate $\hat{x}$ to lie on the arc $\hat{A}\hat{B}$ (see Figure 7.2b), hence $\hat{y}$ (see Figure 7.2a) is the point closest to $\hat{x}$ that lies on the arc $\hat{A}\hat{B}$. The third algorithm is the classical HIO method without any alterations. In fact, we tested also a phase-aware modification of the GS algorithm, however, its results were consistently worse than those of PA-HIO, so we omit them. The current modification of our method uses only one stage. That is, we abandoned the first stage that was used in the original method from Chapter 5 to find a feasible $x$ by solving the convex problem in the original algorithm. This stage is no longer necessarily because the phase bounds in the Fourier domain are the main reasons for success. The additional restrictions on $|\hat{x}|$ used in the original method can be viewed as heuristic constraint added (with smaller weight) for the reasons given in the discussion that follows Equation (6.10). This also made the choice of the initial $x$ straightforward: $x^0 = 0$.

We first demonstrate how the phase uncertainty interval affects our ability to reconstruct the images. A set of 51 uncertainty intervals was chosen in the range from zero to 2.5 radians. For each interval of uncertainty its endpoints were chosen at random so that the true phase was uniformly distributed inside it. We ran our quasi-Newton optimization algorithm one hundred times (each time generating new phase bounds), checking at each run whether it was successful or not. A run was considered successful if the error in the Fourier domain (as defined by Equation (6.1)) was below $0.5 \times 10^{-6}$ after 1000 iterations. As is evident from Figure 6.3 the reconstruction always succeeded as long as the phase uncertainty was below $\pi/2$, in perfect agreement with our analysis. It is also evident that, for images with tight support information, successful reconstruction can be expected even for significantly
rougher phase estimation. Moreover, the algorithm converges very fast and the above threshold is usually reached after 250-300 iterations for the loose-support image and only 80 iterations for the image with tight support as is apparent from Figure 6.5 and 6.6.

![Figure 6.3: Reconstruction success rate of our method as a function of phase uncertainty interval.](image)

Next, we demonstrate in detail the reconstruction results in the case where the Fourier phase uncertainty interval is 1.57 radians. Note that without the phase bounds the HIO method cannot reconstruct images with loose support. Images with tight support information are usually reconstructed successfully, though they may undergo some trivial transformation, for example, axis reversal. As is evident from Figure 6.4 our methods (quasi-Newton and PA-HIO) produce very good visual quality. HIO, on the other hand has problems with the loose-support image (as expected) but the second case seems to yield acceptable quality. However, visual assessment does not provide full insight into the quality of reconstruction and tells nothing about its speed. Quantitative results are given in Figure 6.5 and 6.6, from which it is evident that our quasi-Newton method significantly outperforms HIO and PA-HIO in terms of speed.
Figure 6.4: Reconstruction results with phase uncertainty interval of 1.57 radians. 
Upper row—loose support: (a) our method (after 250 iterations), (b) PA-HIO (after 1000 iterations), and (c) HIO (after 1000 iterations). 
Lower row—tight support: (d) our method (after 80 iterations), (e) PA-HIO (after 1000 iterations), and (f) HIO (after 1000 iterations).
It is important to point out two things before evaluating the quantitative results presented in Figures 6.5, and 6.6. First, all presented methods are iterative by nature and every one of them uses two Fourier transforms per iteration. Hence, comparing the number of iterations is well justified and gives a good estimation of the reconstruction speed because the Fourier transforms constitute the major computational burden. Second, it is obvious that images with loose support lead to non-unique solutions. Hence, a small value of the objective function does not necessarily mean small error in the object domain. This explains the results in Figure 6.5. Another important observation is that the HIO and PA-HIO methods do not enforce the off-support areas (padding) to be zero. Hence, these methods may give a large error in the Fourier domain, while the error in the object domain (after we discard the off-support parts) may be small. This phenomena is also evident in Figure 6.5, and 6.6.

Figure 6.5: Error behavior in the case of loose support: (a) Fourier domain error $\|F x - | \hat{z} \|^2$, (b) object domain error $\| x - | z \|^2$. 
6.3 Concluding remarks

In this chapter we presented a new analysis explaining why continuous optimization methods fail when applied to the phase retrieval problem. On the basis of this observation we gave a heuristic explanation why local minima of a functional associated with the phase retrieval problem can be expected to be global ones in the situation where the Fourier phase error does not surpass $\pi/2$ radians. This, in turn, opens the door for continuous optimization methods whose rate of convergence and ability to incorporate additional information in the computational scheme significantly exceeds those of HIO. We also present such an algorithm and demonstrate that its reconstruction speed is significantly faster than that of HIO, even when the phase constraints are also employed in the latter.

The analysis and the methods developed in this chapter are used in the next chapter to perform phase retrieval in situations where the Fourier phase uncertainty is greater than $\pi/2$ radians by using a type of bootstrapping approach.

Figure 6.6: Error behavior in the case of tight support: (a) Fourier domain error: $\| |Fx| - |\hat{z}| \|^2$, (b) object domain error: $\| x - z \|^2$. 
7 Phase retrieval combined with digital holography*

In this chapter we take our algorithm developed in the previous chapters into a new niche: signal reconstruction from two intensity measurements made in the Fourier plane. One is the Fourier magnitude of the sought image, as in classical phase retrieval, and the second is the intensity pattern resulting from the interference of the original signal with a known reference beam, as in the Fourier domain holography. Although either one of these measurements may, in theory, be sufficient for successful reconstruction of the unknown image, our method provides significant advantages over such reconstructions. For example, comparing with reconstruction from the Fourier magnitude alone by HIO, our method gives a much faster speed and better quality in case of noisy measurements as we showed earlier. Furthermore, unlike classical holography methods, our algorithm does not require any special design of the reference beam. Finally and most importantly, very good reconstruction quality is obtained even when the reference beam contains severe errors.

7.1 Basic reconstruction algorithm

Let us start with the notation used throughout the chapter. The unknown two-dimensional signal that we wish to reconstruct is represented by the complex-valued function \( z(\xi, \eta) = |z(\xi, \eta)| \exp[j\varphi(\xi, \eta)] \). To address the phase of a complex-valued number we use the angle notation, as before: \( \angle z(\xi, \eta) \equiv \varphi(\xi, \eta) \). Our measurements are done in the Fourier plane \((\xi', \eta')\), hence the transformation that \( z \) undergoes when transforming from the \((\xi, \eta)\) plane to the \((\xi', \eta')\) plane is simply the unitary Fourier transform

\[
\hat{z}(\xi', \eta') = \mathcal{F}\{z(\xi, \eta)\}.
\]

(7.1)

Hereinafter, we use the usual convention that a pair of symbols like \( x \) and \( \hat{x} \) denotes a signal in the \((\xi, \eta)\) plane (also known as the object domain) and its counterpart in the \((\xi', \eta')\) plane (also referred to as the Fourier domain), respectively. For the sake of brevity, we may omit the location designator \((\xi, \eta)\) or \((\xi', \eta')\) and use \( x \) or \( \hat{x} \) when the entire signal is considered.

The main purpose of our work is to develop a robust reconstruction method that can tolerate severe errors in the reference beam. To this end we use the reference beam only for estimating the Fourier phase of the sought image. Once a rough phase

*The material presented in this section is currently in preparation for submission to a journal.
estimate is available we can use the method of phase retrieval with approximately known Fourier phase that was developed in the previous chapters.

The two measurements available at our disposal are used as follows. \( I_1 \) provides the Fourier magnitude of the sought image via the simple relationship between the two:

\[
I_1(\xi', \eta') = |\hat{z}(\xi', \eta')|^2.
\]  

The second measurement reads

\[
I_2(\xi', \eta') = |\hat{z}(\xi', \eta') + \hat{R}(\xi', \eta')|^2,
\]  

where \( \hat{R}(\xi, \eta) \) denotes a known reference beam that is used to obtain the Fourier phase estimate as described below. One possible schematic setup that provides these measurements is shown in the next figure.

Note that \( \hat{R} \) is not necessarily a Fourier transform of some physical signal \( R \) in the object domain. This means that \( \hat{R} \) can be formed directly in the Fourier plane without forming first \( R \) and applying then an optical Fourier transform to obtain \( \hat{R} \). Nevertheless, there exists the mathematical inverse

\[
R(\xi, \eta) = \mathcal{F}^{-1}\{\hat{R}(\xi', \eta')\}.
\]  

\[Figure 7.1: \text{Schematic representation of the experiment.}\]
whose properties, such as extent, magnitude, etc. can be considered. The only requirement of \( \hat{R} \) is that it must not vanish in the region of our measurements. This is an important point that provides an advantage for our method over the classical holography techniques. We shall elaborate more on this in Section 7.2.

Let us now describe how \( \hat{z} \)'s phase information is extracted from \( I_2 \), and, more importantly, how it is used in our reconstruction method. Consider the two signals:

\[
\hat{z}(\xi', \eta') = |\hat{z}(\xi', \eta')| \exp[j\phi(\xi', \eta')] ,
\]

and

\[
\hat{R}(\xi', \eta') = |\hat{R}(\xi', \eta')| \exp[j\psi(\xi', \eta')] .
\]

The intensity pattern of their interference can be written as:

\[
I_2(\xi', \eta') = |\hat{z}(\xi', \eta') + \hat{R}(\xi', \eta')|^2
= |\hat{z}(\xi', \eta')|^2 + |\hat{R}(\xi', \eta')|^2 + \hat{z}^*(\xi', \eta') \hat{R}(\xi', \eta') + \hat{z}(\xi', \eta') \hat{R}^*(\xi', \eta')
= |\hat{z}(\xi', \eta')|^2 + |\hat{R}(\xi', \eta')|^2 + 2|\hat{z}(\xi', \eta')| |\hat{R}(\xi', \eta')| \cos[\phi(\xi', \eta') - \psi(\xi', \eta')] .
\]

From this formula we can easily extract the difference between the unknown phase \( \phi(\xi', \eta') \) and the known phase \( \psi(\xi', \eta') \):

\[
\cos[\phi(\xi', \eta') - \psi(\xi', \eta')] = \frac{I_2(\xi', \eta') - |\hat{z}(\xi', \eta')|^2 - |\hat{R}(\xi', \eta')|^2}{2|\hat{z}(\xi', \eta')| |\hat{R}(\xi', \eta')|} .
\]

This gives us:

\[
\phi(\xi', \eta') = \psi(\xi', \eta') \pm \alpha(\xi', \eta') ,
\]

where

\[
\alpha(\xi', \eta') = \arccos \left[ \frac{I_2(\xi', \eta') - |\hat{z}(\xi', \eta')|^2 - |\hat{R}(\xi', \eta')|^2}{2|\hat{z}(\xi', \eta')| |\hat{R}(\xi', \eta')|} \right] .
\]

This expression is well defined, as \( |\hat{R}(\xi', \eta')| \) is assumed to be non-zero everywhere in the region of interest, and the places where \( |\hat{z}(\xi', \eta')| = 0 \) can be simply excluded from our consideration as there is nothing to be recovered because their phase has no influence. We assume that \( \pm \alpha \), that is, the difference between the phases \( \phi \) and \( \psi \), lies within the interval \( [-\pi, \pi] \), hence, no phase unwrapping is necessary. The phase \( \phi(\xi', \eta') \) can assume either one (rarely) or two possible values at every location. The two possible situations are shown in Figure 7.2.
Figure 7.2: Given a reference beam (black) whose magnitude and phase are known, and an unknown signal of known magnitude (dotted circle radius), one can try to find the phase of the unknown signal by measuring the magnitude of the sum (solid circle radius). Evidently, in most cases there are two possible solutions (a). However, in certain cases, the is only one solution (b).

Hence, if the intensity $I_2(\xi', \eta')$ is sampled at $N$ points there are generally $2^N$ possible solutions $\hat{z}(\xi', \eta')$ and, consequently, the same number of possible reconstructions $z(\xi, \eta)$. (Here we consider the worst case scenario where all sampled values give rise to two solutions.) To guarantee a unique and meaningful reconstruction we must use additional information about the sought signal $z(\xi, \eta)$. In the phase retrieval problem, as well as in holography, it is usually assumed that $z(\xi, \eta)$ has limited support, namely, part of the image is occupied by zeros. In practice, it is usually assumed that, in each direction, half or more pixels of $z(\xi, \eta)$ are zeros. To capture this information in the Fourier domain one should “over-sample” by a factor of two (or more) in each dimension. Hence, if the known (not necessarily tight) support area of $z$ is $n \times n$ pixels, then in the Fourier plane it must be sampled with a sensor of size $2n \times 2n$ pixels. Such “over-sampling” usually guarantees unique (up to trivial transformations: shifts, constant phase factor, and axis reversal) reconstruction in the case of the classical phase retrieval problem, where only $|\hat{z}|$ is available (Hayes, 1982). It is not known whether this two-fold oversampling is absolutely necessary in our case where two measurements are available. However, our experiments indicate that for a general complex-valued signal it still seems to be necessary to over-sample by a factor of two. Hence, the reconstruction problem reads: find $z(\xi, \eta)$ such that $|\hat{z}|$ is known, $\angle(\hat{z}) = \psi \pm \alpha$, and $z(x_O, y_O) = 0$. Here, $\alpha$ is the known phase difference between $\hat{z}$ and $\hat{R}$ as defined by Equations (7.9) and (7.10); $(x_O, y_O)$ denotes the known off-support area in the $(\xi, \eta)$ plane.

The problem is combinatorial in nature, and many different methods can be applied
to find a solution. Our method is based on replacing the equality \( \angle(\hat{z}) = \psi \pm \alpha \) with the less strict inequality
\[
\psi - \alpha \leq \angle(\hat{z}) \leq \psi + \alpha .
\] (7.11)
By this relaxation we reduce the original problem into the phase retrieval problem with approximately known phase. For this situation we have developed an efficient Quasi-Newton optimization method based on convex relaxation. Note that the problem we are facing here:
\[
\begin{aligned}
\min_x & \quad \| |\mathcal{F}\{x\}| - |\hat{z}| \|^2 \\
\text{subject to} & \quad \psi - \alpha \leq \angle(\mathcal{F}\{x\}) \leq \psi + \alpha , \\
& \quad x(x_O, y_O) = 0 ,
\end{aligned}
\] (7.12)
is exactly the same as the one we solved in Chapters 5, and 6. Hence, we use the same convex relaxation as we did before. Likewise, the solution of the above minimization problem \( x \) is not guaranteed to be equal to the sought signal \( z \). Due to the relaxation we performed, the phase of \( \hat{x} \) is allowed to assume the continuum of values in the interval \([\psi - \alpha, \psi + \alpha]\) instead of the two discrete values \( \psi \pm \alpha \). However, due to the uniqueness of the phase retrieval problem, the phase of \( x \) may differ from the phase of \( z \) only by a constant. That is, \( x(\xi, \eta) = z(\xi, \eta) \exp[jc] \) for some scalar \( c \) (see (Hayes, 1982) for details). This does not pose any problems, as only the relative phase distribution inside the support area of \( z(\xi, \eta) \) is usually of interest. Moreover, in the case where the absolute phase is required, \( c \) can be recovered by adding a post-processing step that solves the one-dimensional optimization problem:
\[
\min_c \quad \| |\hat{x} \exp[jc] + \hat{R}| - I_2^{1/2} \|^2 .
\] (7.13)
Note that we intentionally do not add a penalty term like \( \| |\hat{x} + \hat{R}| - I_2^{1/2} \|^2 \) into the main minimization scheme as defined by Equation (7.12). Adding such a term would introduce a strong connection between \( x \) and the reference beam \( \hat{R} \). This connection will inevitably deteriorate the quality of reconstruction when the reference beam \( \hat{R} \) contains errors.

There is, however, an important dissimilarity between the current situation and the one we considered in the previous chapter: the phase uncertainty interval can be as large as \( \pi \) radians. Nevertheless, our experiments indicate that the reconstruction is stable and its speed is very fast (see Figure 7.6). Moreover, it can be further accelerated, and our experiments indicate that more aggressive oversampling (zero padding in the object domain) results in faster convergence in terms of the number of iterations. In fact, use of a special reference beam can, in theory, result in a trivial non-iterative reconstruction in a way similar to holography. However, such special reference beams may not be easily realizable in physical systems and the quality of the reconstructed signal is strongly influenced by the quality of the reference beam. We discuss this setup in the next section and compare its sensitivity to possible errors in \( \hat{R} \) against our method in Section 7.4.
7.2 Relation to holography

Our method was initially developed for the phase retrieval problem. However, the use of interference patterns creates a strong connection with holography. Therefore, it may be pertinent to discuss the advantages that our method provides over the classical holographic reconstruction. Note that our discussion is limited to basic holography only, and no attempt is made to cover all possible setups and techniques that can be used in digital holography. We nonetheless believe that this novel approach can compete with or improve upon existing algorithms used in digital holography.

In classical holography one uses a specially designed reference beam so as to allow easy non-iterative recovery of the sought image. This has an obvious advantage over iterative methods, especially when the speed of the reconstruction is of high importance. However, reliance on the reference beam means that reconstruction quality may deteriorate badly when the reference beam contains errors, that is, when it differs from the “known” values. To review the non-iterative reconstruction process we denote convolution. Note that in this case the fourth and the third terms are equal to the sought wavefront $z(\xi,\eta)$, and its Hermitian counterpart $z^*(-x,-y)$ convolved with $A(\xi,\eta) \otimes R(\xi,\eta)$ and $A(\xi,\eta) \otimes R^*(-x,-y)$, respectively. The best possible choice is $A(\xi,\eta) = \delta(\xi,\eta)$ and $R(\xi,\eta) = \delta(\xi - \xi_0, \eta - \eta_0)$, where $\delta(\xi,\eta)$ is the Dirac delta function. In this case the obtained wave becomes:

$$B(\xi,\eta) = z(\xi,\eta) \otimes z^*(-\xi,-\eta) + \delta(\xi,\eta) + z^*(-\xi+\xi_0,-\eta+\eta_0) + z(\xi-\xi_0,\eta-\eta_0).$$  \hspace{1cm} (7.15)$$

Hence, if $\xi_0$ and/or $\eta_0$ are large enough, the four terms in the above sum will not overlap in the $(\xi,\eta)$ plane. Thus, we can easily obtain the sought signal $z(\xi,\eta)$, albeit shifted by $(\xi_0,\eta_0)$, provided that the spatial extent of $z(\xi,\eta)$ is limited by the box $\xi \in [-L_\xi, L_\xi], \eta \in [-L_\eta, L_\eta]$. The spatial extent of the autocorrelation $z(\xi,\eta) \otimes z^*(-\xi,-\eta)$ is twice as large, that is, limited by the box $\xi \in [-2L_\xi, 2L_\xi], \eta \in [-2L_\eta, 2L_\eta]$. Hence, to avoid overlapping we must have $\xi_0 > 3L_\xi$, or $\eta_0 > 3L_\eta$. 

$$\text{71}$$
Thus, in theory, one can generate an ideal delta function in the $(\xi, \eta)$ plane located at a sufficient distance from the support area of $z(\xi, \eta)$. In the Fourier domain, this delta function corresponds to a plane wave arriving at a certain angle at the plane of measurements. If such a construction is possible, then a simple inverse transform of the intensity obtained in the Fourier plane is sufficient to obtain the sought signal $z(\xi, \eta)$. However, as mentioned earlier, this approach has some drawbacks. First, it is impossible to create an ideal delta function. Any physical realization will necessarily have a finite spatial extent, and this will result in a “blurred” reconstructed image. Note that the term “blurring” describes well the resulting image in the case where $z(\xi, \eta)$ is real-valued or has constant phase. In the more general case, where the phase of $z(\xi, \eta)$ varies at non-negligible speed, the result appears more distorted (see Figure 7.4). The second drawback is the sensitivity of this method to errors in $\hat{R}$. In Section 7.4 we demonstrate how the quality of reconstruction depends on the error in $\hat{R}$ (see Figures 7.8, 7.9, and 7.10). Our method, on the other hand shows very little sensitivity to the reference beam shape. Moreover, its modification described in the next section allows the reference beam to contain severe errors without deteriorating significantly the quality of reconstruction.

### 7.3 Reconstruction method for imprecise reference beam

Here we consider the situation where the reference beam is not known precisely, that is, we assume that the phase of $\hat{R}$ contains some unknown error. It is easy to verify that if the reference beam phase $\psi(\xi', \eta')$ has error $\epsilon(\xi', \eta')$ then the sought phase $\phi(\xi', \eta')$ becomes

$$
\phi(\xi', \eta') = \psi(\xi', \eta') + \epsilon(\xi', \eta') \pm \alpha(\xi', \eta'),
$$

in a manner similar to Equation (7.9). We do not consider errors in the magnitude $|\hat{R}|$ for several reasons. Many aberrations manifest themselves through phase distortion (Goodman, 2005). Also, the magnitude of $\hat{R}$ can be measured. Moreover, looking at the above equation, it is obvious that any error in $\psi$ can be viewed as an error in $\alpha$. That is, the situation would be the same if the reference beam phase $\psi$ were known precisely, while the difference $\alpha$ would contain some errors. This observation is relevant because errors in the phase $\alpha$ can arise from many different sources, including imperfect measurements and errors in the reference beam magnitude.

The true error $\epsilon(\xi', \eta')$ is, of course, unknown. Hence, we assume just an upper bound (assumed known) on the absolute phase error:

$$
\psi - \epsilon - \alpha \leq \angle(\hat{z}) \leq \psi + \epsilon + \alpha,
$$

as in Equation (7.11). This time, however, the phase uncertainty interval may be larger than $\pi$ radians which makes our method inapplicable. On the other hand, limiting the phase uncertainty interval by $\pi$ radians will prevent us from reconstructing the precise
image, because the true phase may lie outside this interval. A possible solution is to measure the intensity of the reference beam and then to reconstruct its phase using the method presented in Chapter 6, because this problem itself can be seen as a phase retrieval with approximately known phase. However, taking another measurement may be undesirable, and therefore we developed the following reconstruction method:

**Step 1:** Set the phase uncertainty interval as defined by Equation (7.11) (as if there were no errors in the reference beam phase).

**Step 2:** Solve the resulting minimization problem, obtaining a solution $x(\xi, \eta)$.

**Step 3:** If not converged, set the phase uncertainty interval to $[\angle(\hat{x}) - \pi/2, \angle(\hat{x}) + \pi/2]$. Clip it, if applicable, to the limits defined by Equation (7.17) and go to Step 2.

In this algorithm we perform a number of outer iterations, each time updating the phase uncertainty interval. This approach leads to a successful reconstruction method even in cases where the reference beam contains severe errors. The results are much better than those of non-iterative holographic reconstruction (see Figures 7.8, and 7.9). This improvement is achieved by decoupling the reconstruction problem (which becomes the pure phase retrieval with approximately known phase) and the erroneous interferometric measurements.

### 7.4 Experimental results

The method was tested on a variety of images with similar results. Here we present numerical experiments conducted on one natural image so as to allow easy perception of the reconstruction quality under various conditions. The image intensity (squared magnitude) is shown in Figure 7.3.

![Figure 7.3: Original image (intensity).](image)

The technical details of the image are as follows: the size is $128 \times 128$ pixels, and pixel values (amplitude) vary from 0.2915 to 0.9634 on a scale of 0 to 1, with mean value of 0.6835. These parameters will become important later, when we will consider the reference beam design, and when we will assess the reconstruction quality. Since
the original image is a photograph, it does not have any phase information. Hence, we generated three different phase distributions to account for the assortment of possible real-world problems where our method can be applied. The first distribution assumes that the image is non-negative real-valued, that is, the phase is zero everywhere. The second distribution is designed to mimic a relatively smooth phase. To this end, the phase is set to be proportional to the image values (scaled to the interval \([-\pi, \pi]\)). Finally, in the third distribution the phase is chosen at random, uniformly spread over the interval \([-\pi, \pi]\). This distribution is designed to show the behavior of our reconstruction method in cases where the true phase varies rapidly. We also consider three possible reference beams, again, to demonstrate the robustness of our method. The first reference beam is an ideal delta-function in the \((\xi, \eta)\) plane, located at the coordinate \((256, 256)\) so that the holographic condition is satisfied. With this reference beam exact reconstruction is obtained as long as the sampling in the Fourier domain is sufficiently dense \((512 \times 512\) pixels, or more). We do not present the visual results of reconstruction for this reference beam as both methods produce images that are indistinguishable from the true image. The speed of convergence of our method is shown in Figure 7.6. Later, we show also how the reconstruction quality of both methods is affected by Fourier phase errors in the reference beam. Before this, we demonstrate the effect of departure from the ideal delta function: the second reference beam is a small square of size \(3 \times 3\) pixels, located at the coordinates \((256 : 258, 256 : 258)\). In this setup the reconstruction quality of the holographic method is degraded, as evident from Figure 7.4.
Figure 7.4: Image (intensity) reconstructed by the holographic technique using a small square as the reference beam: (a) the image is real-valued, (b) image phase varies slowly, (c) image phase is random (varies rapidly).

It is also evident that faster variations in the object phase result in greater deterioration in the reconstruction, in agreement with our expectations. Our method, on the other hand, is insensitive to the reference beam form. In Figure 7.5 we demonstrate our reconstruction results for the aforementioned small square as the reference beam (the first row), and for another reference beam that was formed in the Fourier plane by combining unit magnitude with random phase (in the interval $[-\pi, \pi]$). This beam, of course, is not suitable for holography, as its extent in the object plane occupies the whole space.
Figure 7.5: Image reconstructed (intensity) by our method: (a), (b), and (c) — reference beam is a small square and object phase is zero (a), smooth (b), random (c). (d), (e), and (f) — reference beam is random and object phase is zero (d), smooth (e), random (f).
Reconstruction is very fast and, in fact, is almost independent of the sought image and reference beam type. Figure 7.6 demonstrates that less than 20 iterations are required to solve the minimization problem as defined by Equation (7.12).

Figure 7.6: Reconstruction speed of our method: (a) real valued image, (b) image phase is smooth, (c) image phase is random.

In all these examples we assume perfect knowledge of the reference beam. Next we consider the situation where the actual reference beam does not match the expected signal in the Fourier plane. Following our discussion in Section 7.3, we evaluate how the reconstruction quality of the holographic approach and our method are affected by errors in the reference beam Fourier phase. We use again the three aforementioned models of the sought image (real-valued, smooth phase, and random phase) and the three reference beams (delta function, small square, and random). From Figure 7.7 it is evident that in all these cases we were able to solve the minimization problem to sufficient accuracy as long as the phase error was below 25%. That is, our method can tolerate reference beam Fourier phase errors of up to \( \pi/2 \) radians. The sharp discontinuity that happens at this value has a simple explanation: phase error greater
than $\pi/2$ radians can result in the phase uncertainty interval greater than $2\pi$ radians (see Equation (7.17)). Hence, all phase information is lost.

Figure 7.7: Fourier domain ($||\hat{x}|-|\hat{z}||^2$) error vs. phase error in the reference beam: (a) real valued image, (b) image phase is smooth, (c) image phase is random.

A comparison with the holographic reconstruction is given in Figure 7.8 where the error norm in the object domain is depicted. Note that the objective function values are about $10^{-10}$, hence, one would expect the object domain error norm to be of order $10^{-5}$ (the difference stems from the fact that the objective function uses squared norm). This is not so in the case of complex-valued images and the random reference beam. This effect is due to the relaxation we perform in the Fourier phase, as discussed in Section 7.1. It does not change the relative phase distribution, but all phases can get a constant addition. This can be corrected by solving the one dimensional minimization defined by Equation (7.13). Figure 7.9 depicts the corrected values yielded by this process. Visual results comparing the holographic reconstruction with our method are provided in Figure 7.10.
Figure 7.8: Object domain error ($\|x - z\|$) vs. phase error in the reference beam: (a) real valued image, (b) image phase is smooth, (c) image phase is random.
Figure 7.9: Corrected object domain error (\(|x - z|\)) vs. phase error in the reference beam: (a) real valued image, (b) image phase is smooth, (c) image phase is random.
Figure 7.10: Image (intensity) reconstructed by the holographic method (upper row) and our method (lower row). Object phase is random, and the reference beam is a delta function with Fourier phase errors: (a), (b), and (c) — phase error is 1%, 10%, and 20% respectively (d), (e), and (f) — phase error is 1%, 10%, and 20% respectively.
As the results show, our method demonstrates a substantial advantage over ordinary holographic reconstruction. It is remarkable that even when the minimization is not particularly successful (in cases of very large phase errors) our reconstruction is still closer to the true signal than the holographic method. This success is due to our approach of decoupling the phase retrieval from the interferometric measurements. As mentioned earlier, we deliberately avoid strong dependence on the reference beam. The interference pattern is only used to estimate the Fourier phase bounds. The results indicate that this approach is well justified.

### 7.5 Concluding remarks

In this chapter we presented a new reconstruction method from two intensities measured in the Fourier plane. One is the magnitude of the sought signal’s Fourier transform, and the other is the intensity resulting from the superimposition of the original image and an approximately known reference beam. While the method was originally developed for the phase retrieval problem, it can be useful in digital holography, because it poses less stringent requirements on the reference beam. The method is designed specifically to allow severe errors in the reference beam without compromising the quality of reconstruction. Numerical simulations justify our approach, exhibiting reconstruction that is superior to that of holographic techniques.

It is also important to note that this method can, potentially, be used in the classical phase retrieval problem without knowing the Fourier phase withing the error of $\pi/2$ radians as was done in Chapters 5, and 6 as it uses some sort of bootstrapping technique that allows much rougher phase estimation.
8 Designing boundaries for phase retrieval

In this chapter we develop an algorithm for signal reconstruction from the magnitude of its Fourier transform in a situation where some (non-zero) parts of the sought signal are known. Although our method does not assume that the known part comprises the boundary of the sought signal, this is often the case in microscopy: a specimen is placed inside a known mask, which can be thought of as a known light source that surrounds the unknown signal. Therefore, in the past, several algorithms were suggested that solve the phase retrieval problem assuming known boundary values (Hayes & Quatieri, 1982; Hayes & Quatieri, 1983; Fienup, 1983; Fienup, 1986). Unlike our method, these methods do rely on the fact that the known part is on the boundary.

Besides the reconstruction method we give an explanation of the phenomena observed in previous work: the reconstruction is much faster when there is more energy concentrated in the known part (Fiddy et al., 1983; Fienup, 1986). Quite surprisingly, this can be explained using our previous results on phase retrieval with approximately known Fourier phase.

8.1 Review of existing methods

Of course, it is possible to use the methods reviewed/developed in the previous sections. One can apply the HIO algorithm and hope not to be entrapped in a situation where the method stagnates. However, this approach is not optimal, as it does not use the additional information available in this case. Note that we cannot use the reconstruction method developed in the previous chapter because in the current setup we assume that only one measurement is available. Our method will be presented later. At the time being let us start by reviewing other methods suggested in literature.

In accordance with the convention adopted in the previous chapters, the unknown signal is denoted by $z(m, n)^\dagger$. The signal is assumed to have a finite support, specifically, it vanishes outside the box $[0, M - 1] \times [0, N - 1]$, and our goal is, as before, to reconstruct it from the (oversampled) magnitude of its Fourier transform $|\hat{z}(p, q)|$, where

$$\hat{\epsilon}(p, q) = |\hat{\epsilon}(p, q)| \exp(j\phi(p, q)) \equiv \mathcal{F}[z(m, n)] \quad .$$ *(8.1)*

*The material presented in this section is currently in preparation for submission to a journal.

†Here we explicitly assume that the signal is discrete.
In our implementation, $F$ the unitary Fourier transform. This regularization constant is chosen with only one purpose: to make the distance (norm) in the Fourier domain equal that in the object domain. Thus, by the discrepancy (error) in the Fourier domain one can easily estimate the error in the object domain. Hence,

$$F[z(m,n)] = (PQ)^{-1/2} \sum_{m=0}^{P-1} \sum_{n=0}^{Q-1} z(m,n) \exp[-j2\pi(mp/P + nq/Q)], \quad (8.2)$$

where $m, p = 0, 1, \ldots, P - 1$ and $n, q = 0, 1, \ldots, Q - 1$. In addition, we require the “two-fold oversampling” in the Fourier domain, that is $P = 2M - 2$, $Q = 2N - 2$. The purpose of this oversampling is to capture the information that $z(m,n)$ has only limited support. With these definitions, there exists one well-known relation between the squared magnitude of the Fourier transform and the linear (as opposed to cyclic) auto-correlation (denoted by $\star$) function of $z(m,n)$:

$$r(i, j) \equiv z(m,n) \star z(m,n) \equiv \sum_{m=0}^{M-1} \sum_{n=0}^{N-1} z(m,n)z^*(m-i,n-j) \quad (8.3)$$

$$= F^{-1}(|\hat{z}(p,q)|^2),$$

where $r(i,j)$ is of size $(2M-1) \times (2N-1)$ pixels defined over the region $1 - M \leq i \leq M - 1$, $1 - N \leq j \leq N - 1$. This relation between the magnitude of the Fourier transform and the auto-correlation function of the sought signal was used by Hayes and Quatieri to develop an elegant algorithm for finding $x^*$ when its boundaries are known (Hayes & Quatieri, 1983; Hayes & Quatieri, 1982). The algorithm assumes that the boundaries, that is, the first row $z(0,:)^\dagger$, the last row $z(M-1,:)$, as well as the first and the last columns: $z(:,1)$, and $z(:,N-1)$, are known. The algorithm is iterative and at each iteration it recovers two new unknown rows (or columns) of $x$. The authors demonstrated that the $k$-th iteration of the algorithm reduces to a simple matrix (pseudo) inverse to solve the following system of equations

$$\begin{bmatrix} F & L \end{bmatrix} \begin{bmatrix} x(N-k,:) \\ x(k,:) \end{bmatrix} = \begin{bmatrix} \hat{r}(N-k,:) \end{bmatrix}, \quad (8.4)$$

where the matrices $F$ and $L$ correspond to the cross-correlation with the first and the last rows, respectively. The right-hand side $\hat{r}(N-k)$ is obtained from the $(N-k)$-th row of the auto-correlation function $r(i,j)$ from which the contribution of already recovered rows $1, 2, \ldots, k-1$ and $M-2, M-3, \ldots, M-k+1$ has been subtracted. For a more detailed description we address the reader to the original articles. The most appealing property of this algorithm is that it requires only a small, known

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*Again, $x$ is not necessarily equal to $z$ due to possible non-uniqueness.

†Here we use MATLAB notation, where a colon (:) denotes the entire vector of indexes of the corresponding dimension.
in advance, number of iterations until the whole signal \(x(m, n)\) is recovered. The authors also provided conditions that they claimed were sufficient to guarantee uniqueness of the reconstruction. The latter, however, were proven to be incorrect (see (Fienup, 1986)). Nonetheless, the biggest problem with this algorithm is not the non-uniqueness of reconstruction, because, as we already mentioned, the two-dimensional phase retrieval solution is usually unique (up to trivial transformations) to start with. The main difficulty that makes the algorithm impractical for all but tiny problems is its numerical instability. It can easily be shown that the error grows exponentially due to the recurrent nature of the algorithm. Even if we assume that the measurements are ideal, containing absolutely no error, each iteration of the algorithm will introduce a small error due to the finite computer precision. In the next iteration, the norm of this error will be increased by a factor proportional to the condition number of the matrix \([F \mid L]\). The new error will be further amplified (by the same factor) in the next iteration, and so on. This will result in extremely fast (exponential) error growth. This is demonstrated in Figure 8.4 where a small (128 \(\times\) 128 pixels) image was reconstructed by the HQ algorithm in the horizontal direction, that is, reconstructing column after column. This exponential growth is observed whenever the condition number of the matrix \([F \mid L]\) is greater than one. It can equal unity in some very special cases, for example, when the known boundaries contain a single delta function. This situation was considered in (Fienup, 1983; Fienup, 1986), and in (Fiddy et al., 1983), although in the latter it was not used directly for the reconstruction—the authors added this condition to guarantee the uniqueness of the reconstruction. Moreover, all of them observed empirically that the reconstruction was faster when the known part contained more energy. This observation is common, even though the authors use different reconstruction methods. None of them, however, provided an explanation for this phenomenon. In the next section we present our reconstruction routine and explain why a “strong” known part leads to a fast and stable reconstruction. Additionally, we will consider the influence of noise in the measurement—another issue that has been largely overlooked in previous works despite its enormous importance.

8.2 Our reconstruction method

First, we must consider the source of the known boundary. In microscopy, it is often natural to create a mask (for transparencies) or a bed (for light reflecting objects) whose boundaries are known and designed in a way that leads to easy image reconstruction. An example of such a mask is shown in Figure 8.1.
The object here is assumed to be transparent, so the white areas correspond to simple windows in an opaque material (shown in black). The big window in the center is where the object is placed. The whole construction is then illuminated by a coherent plane wave such that the small windows in the mask can be assumed to be of known intensity. There is nothing special about the plane wave here, the most important point is that part of the image is known. This setup allows us to formulate the following minimization problem to find the unknown signal $x(m,n)$:

$$
\min_x \| |\mathcal{F}[x + b]| - r \|^2 \\
\text{subject to } x(m \in \mathcal{O}_m, n \in \mathcal{O}_n) = 0,
$$

(8.5)

where $r$ denotes the measured Fourier magnitude of the entire signal, $b$ represents the known part (boundary) and $(\mathcal{O}_m, \mathcal{O}_n)$ designate the location of the off-support parts of $x$ (basically, these are the locations occupied by the mask except the central window where the object is located). Note that we, again, use $x$ to denote the reconstruction result because, in general, it may not be equal the sought signal $z$.

Following the discussion in Chapter 7, the mask can, in principle, be constructed in a way that makes the reconstruction trivial: it is sufficient to place only one infinitesimally small window (a delta function) at a sufficient distance from the object. In this case the mask would satisfy the holography conditions and the reconstruction could be as easy as applying a single Fourier transform. However, as was mentioned before, generating a delta function is not possible in practice. Practical mask design must balance between the production costs/difficulties and how helpful it is for the reconstruction process. Hence, here we use simple square windows of relatively large size, located close to the object. Hence, a non-iterative
reconstruction is not possible. However, using a quasi-Newton optimization method to solve (8.5) we were able to get good results, as demonstrated in Section 8.3.

Before we proceed to the numerical results, it is pertinent to discuss briefly the design of the mask. It is not known at the moment what is the best way to design the mask so that the reconstruction would be fast and robust. Our experience shows that the mask should have a strong “presence” in all frequencies. More precisely, we can explain why this situation leads to a fast reconstruction. Consider a single element in the Fourier space. The contribution of the known part (denoted by \( \hat{b}_i \)) is fully determined, that is, its magnitude and phase are known. Hence, the full Fourier domain data at this location is the sum \( \hat{x}_i + \hat{b}_i \) that is located somewhere on the red circle shown in Figure 8.2. Note that the phase \( \angle(\hat{b}_i + \hat{x}_i) \) is known up to \( \pi/2 \) radians when \( \alpha \leq \pi/4 \) (see Figure 8.2). That is, when

\[
\arcsin \left( \frac{|\hat{x}_i|}{|\hat{b}_i|} \right) \leq \frac{\pi}{4} \iff |\hat{b}_i| \geq \sqrt{2}|\hat{x}_i|. \tag{8.6}
\]

\[\text{Figure 8.2: “Strong” known part leads to the phase retrieval problem with approximately known Fourier phase.}\]

The relation in Equation (8.6) is quite interesting: if the energy concentrated in the known part is at least twice as large as the energy in the unknown part\(^*\), the problem reduces to phase retrieval with Fourier phase known within the limit of \( \pi/2 \) radians. Hence, according to our results in Chapter 6, any reasonable algorithm will be able to reconstruct the unknown part. Moreover, the larger the ratio \( |\hat{b}_i|/|\hat{x}_i|\),

\[^*\text{The energy must also be distributed “well” in the frequency domain.}\]
the smaller the Fourier phase uncertainty is. However, there is another aspect that is important in practice—in all physical experiments, the measurements inevitably contain some noise. In our case, where the measurements are light intensity, the noise is well approximated by the Poisson distribution. This means that stronger intensity will result in more noise (though, the signal to noise ratio (SNR) usually increases with the intensity growth). Hence, making the known part too strong compared with the sought signal will make the measurement noise more dominant with the respect to the unknown part and will eventually arrive at the level where the reconstruction is not possible. Hence, one should not just increase the intensity of the known part, as this leads to poor reconstruction quality in case of noisy measurements\(^\ast\). The most appealing approach would be to design a mask that would use the limited power in an efficient way. Without any a priori knowledge about the sought signal, it seems that the optimal way would be to create a mask whose Fourier domain power is spread evenly over all frequencies. Note the relation to the holographic reconstruction where a single small window (delta function) is used, because the Fourier domain power of a delta function is the same over all frequencies. Instead of using a delta function we can obtain a very good approximation to the uniform power spectrum if some randomness is added to the mask windows. It can be random shape of the windows or random values/phases across them. Making random shapes may be more difficult than adding a diffuser into a square window. Hence, in Section 8.3 we demonstrate the reconstruction with mask windows of constant intensity and random intensity. As expected, adding randomness improves the reconstruction speed and noise-immunity.

### 8.3 Numerical results

We experimented with several images, however, the results provided below are limited to two images of size 128 × 128 pixels. These were chosen to represent two different classes of objects. One is a natural image “Lena” already used in our previous experiments. Another is the Shepp-Logan phantom. These images are very popular in other fields. “Lena” is a classical benchmark in the image processing community, because it has a lot of features and delicate details. The second image, “phantom”, is often used as a benchmark in MRI related algorithms. However, its piece-wise constant nature can approximate well objects that are often investigated in microscopy, for example, cells. Besides the above differences these two images differ by their support. Lena’s support is tight, that is, it occupies all the space and no shifts are possible. The phantom, on the other hand, has non-tight support. As we saw earlier, this is an important property for phase retrieval algorithms—complex valued images with non-tight support are much more difficult for the current reconstruction methods like HIO. The image intensities (squared magnitude) is shown in Figure 8.3.

\(^\ast\)Here we assume that the noise grows with the signal, as indeed happens with Poissonian noise.
The images were tested in two different scenarios: first, they were assumed to be real-valued and non-negative; second, they were made complex-valued by adding a phase distribution. Here we present the results for the case where the objects’ phases were chosen to be proportional to their intensity (scaled to the interval $[0, 2\pi]$). This choice corresponds to the case where the phase changes in a relatively smooth manner. However, all our experiments indicate that the particular phase distribution has little effect on the reconstruction, except the case where the object can be assumed real non-negative. In this case the non-negativity prior can be used to speed-up the reconstruction and improve its quality in the case of noisy measurements, as is evident from Figures 8.6, and 8.7.

Let us begin with a demonstration of the HQ algorithm results. Following our discussion in Section 8.1 we expect the error to grow exponentially as we progress from the boundaries toward the image center. This expectation is confirmed by the results shown in Figure 8.4. Note that the reconstructed intensity in Figure 8.4b cannot convey the true error because the image storage format clips all values outside the interval $[0, 1]$. The true error is evident from Figure 8.4c.
Figure 8.4: Error growth in the Hayes-Quatieri recursive algorithm: (a) original image, (b) reconstructed by the HQ algorithm, and (c) actual error growth in the HQ algorithm.

In theory, we can use the same boundary conditions as the HQ algorithm, however, our experiments indicate that the optimization routine used in our method is prone to stagnation when the known boundary (or image part in general) carries little energy. This is, of course, in agreement with our discussion in the previous section: when the boundary carries little energy it does not provide enough information about the Fourier phase. This, in turn, causes line-search optimization algorithms to stagnate (see Chapter 6). This stagnation could be addressed by a more sophisticated reconstruction routine similar, for example, to that in Chapter 7 combined with some techniques from global optimization. However, this approach can be expensive in
the terms of computational effort. Moreover, the stagnation can be alleviated by designing a mask that makes the reconstruction much easier. In our approach the emphasis is put on the simplicity of the mask design. Hence, we used the simple mask shown in Figure 8.1. The mask is of size $256 \times 256$ pixels with 11 square windows of size $20 \times 20$ located approximately on a circle of radius 100 pixels (see Figure 8.1). The objects are placed in the middle of the mask where a special window of size $128 \times 128$ pixels is provided. The area outside this special window comprises the known boundary. When placed in the mask, the test images look as shown in Figure 8.5.
Note also that the algorithm based on the minimization problem defined in Equation (1.5) is very naive and does not try to use the “approximately known Fourier phase” in the case were the energy in the known part is sufficiently large. However, our main goal is to show the influence of the mask design. This influence is essentially independent of the algorithm used for reconstruction.

Let us demonstrate how the energy contained in the known part affects the reconstruction. Recall that our expectation is that a mask that has a strong “presence”
in all Fourier frequencies will better suit the reconstruction process in the noise-less case. The results shown in Figures 8.6, and 8.7 fully support this conjecture. These figures present the reconstruction speed (the objective function minimization rate) for three different magnitudes of the mask’s windows values: 5, 10, and 100. The lowest value (5) was chosen so as to be close to the theoretical ratio between the energy in the known and the unknown parts that is required for guaranteed reconstruction, as defined in Equation (8.6). However, as we can see the value of 5, and even the value of 10 did not result in perfect reconstruction. This phenomenon has two reasons: first, doubling the amount of energy in the known part compared to the unknown part is not sufficient—this energy must be distributed properly in the Fourier domain; second, it may be attributed to the simplicity of the chosen reconstruction algorithm. However, the second reason is less likely in view of the following experiment. In an attempt to create a “better” distribution of the mask’s energy in the Fourier domain we added some “randomness” to the mask by modulating (multiplying) the flat values across the mask’s windows with random values in the interval [0, 1]. This step improved the speed of the reconstruction and its robustness, as is evident from Figures 8.8, and 8.9.

![Figure 8.6: Lena's reconstruction speed with flat mask values: (a) real-valued, (b) complex valued.](image-url)
Figure 8.7: Phantom’s reconstruction speed with flat mask values: (a) real-valued, (b) complex valued.

Figure 8.8: Lena’s reconstruction speed with random mask values: (a) real-valued, (b) complex-valued.
Figure 8.9: Phantom’s reconstruction speed with random mask values: (a) real-valued, (b) complex-valued.

Note that now the reconstruction is successful for all mask values and is very fast. However, despite this fast convergence, one must be careful not to put too much energy into the known part. This approach may harm the reconstruction quality of the unknown part when there is noise in the measurements, as we demonstrate next. In these experiments the measurements (intensity values) were contaminated with Poisson noise with different SNR ranging from 10 to 60 decibels. As is evident from Figures 8.10 and 8.11, the more energy is concentrated in the known part the worse is the reconstruction quality of the unknown part. This phenomenon is, of course, expected. The Poisson noise is signal dependent: higher intensity results in more noise. However, the intensity (energy) of the unknown part remains constant, hence the noise becomes more and more significant compared to it. To obtain the best result one would like to design a mask whose power spectrum will correlate well with the power spectrum of the sought signal. Unfortunately, this approach cannot be implemented, because designing such a mask requires a priori knowledge of the sought signal’s Fourier magnitude, which is unavailable in our case. However, it may be a good approach when the Fourier magnitude of the sought signal is known approximately.
Figure 8.10: Lena’s reconstruction quality: (a) real-valued with flat mask value, (b) complex-valued with flat mask value, (c) real-valued with random mask values, (d) complex-valued with random mask values.
Figure 8.11: Phantom’s reconstruction quality: (a) real-valued with flat mask value, (b) complex-valued with flat mask value, (c) real-valued with random mask values, (d) complex-valued with random mask values.

### 8.4 Concluding remarks

In this chapter we considered the problem often met in the Fourier domain holography: signal reconstruction form the Fourier magnitude of the sum of the sought signal and a reference beam. We provided an explanation to the fact observed in practice: why a strong reference beam leads to a faster reconstruction for a variety of reconstruction methods. Based on this explanation we suggested a “good” boundary (reference beam) design. The latter problem (reference beam design) requires more research, as the optimal reference beam must satisfy at least two requirements in the presence of
signal dependent noise. For example, in the case of Poissonian noise (or any other noise model in which the noise level grows with the signal intensity) the optimal reference beam must be simultaneously “strong” (to aid the reconstruction process) and “weak” (to alleviate) the destructive influence of the noise.

In general, when the Fourier magnitude of the sought image is known approximately, the best mask should have the power spectrum that is about two times larger than that of the sought signal (in each frequency). If the power spectrum of the sought images is unknown, the mask should have a strong presence in all frequencies. In this case, it seems that the best design would be based on some “randomness” in the mask values or shapes.
9 Bandwidth extrapolation using sparsity constraints*, †

In this chapter we present our work on the bandwidth extrapolation (super-resolution) problem with application to Coherent Diffracting Imaging (CDI). CDI is an algorithmic imaging technique where intricate features are reconstructed from measurements of the freely-diffracting intensity pattern (Sayre, 1952; Miao et al., 1999; Miao et al., 2001; Quiney, 2010; Chapman & Nugent, 2010). An important goal of such lensless-imaging methods is to study the structure of molecules (including many proteins) that cannot be crystallized (Sandberg et al., 2007; Chapman et al., 2007; Neutze et al., 2000). Clearly, high spatial resolution and very fast measurement are key features for many applications of CDI. Ideally, one would want to perform CDI at the highest possible spatial resolution and in a single-shot measurement—such that the techniques could be applied to imaging at ultra-fast rates. Undoubtedly, such capabilities would give rise to unprecedented possibilities. For example, observing molecules while they dissociate or undergo chemical reactions will considerably expand the knowledge in physics, chemistry and biology. However, the resolution of all current CDI techniques is limited by the diffraction limit, and therefore cannot resolve features smaller than one half the wavelength of the illuminating light (Lipson et al., 2010), which is considered a fundamental limit in diffractive imaging (Abbe, 1873). Moreover, combining CDI with current sub-wavelength imaging techniques would not allow for rapid single-shot measurements that are able to follow ultra-fast dynamics, because such techniques rely on multiple exposures, either through mechanical scanning (e.g., Scanning Near-Field Microscope (Lewis et al., 1984; Betzig et al., 1991), scanning a sub-wavelength “hot spot” (Di Francia, 1952; Lezec et al., 2002; Huang & Zheludev, 2009)), or by using ensemble-averaging over multiple experiments with fluorescent particles (Yildiz et al., 2003; Hell et al., 2009). Here, we present sparsity-based single-shot sub-wavelength resolution in coherent diffraction microscopy: algorithmic reconstruction of sub-wavelength features from far-field intensity patterns of sparse optical objects. We experimentally demonstrate imaging of irregular and ordered arrangements of 100 nm features with illumination wavelength of 532 nm (green light), thereby obtaining resolutions several times better than the diffraction limit. The sparsity-based sub-wavelength imaging concept relies on minimization of the number

*The work presented in this chapter was done in collaboration with Prof. Segev’s group form the Technion Physics Department, Solid State Institute.
†The material presented in this chapter was submitted to the Nature journal. Part of it was presented at the Frontiers in Optics 2011 conference. Also, part of it was submitted to the CLEO 2012 conference.
of degrees of freedom, and operates on a single-shot basis (Gazit et al., 2009; Szameit et al., 2010; Gazit et al., 2010). Hence, it is suitable for capturing a series of ultrafast single-exposure images, and subsequently improving their resolution considerably beyond the diffraction limit. This work paves the way for ultrafast sub-wavelength CDI, via phase retrieval at the sub-wavelength scale. For example, sparsity-based methods could considerably improve the CDI resolution with x-ray free electron laser (Chapman et al., 2011), without hardware modification. Conceptually, sparsity-based methods can enhance the resolution in all imaging systems, optical and non-optical.

9.1 Background information

Improving the resolution in imaging and microscopy has been a driving force in natural sciences for centuries. Fundamentally, the propagation of an electromagnetic field in a linear medium can be fully described through the propagation of its eigen-modes (a complete and orthogonal set of functions which do not exchange power during propagation). In homogeneous, linear and isotropic media, the most convenient set of eigen-modes are simply plane-waves, each characterized by its spatial frequency and associated propagation constant (see (Goodman, 2005, Section 3.10)). However, it is also known that when light of wavelength $\lambda$ propagates in media with refractive index $n$, only spatial frequencies below $n/\lambda$ can propagate, whereas all frequencies above $n/\lambda$ are rendered evanescent and decay exponentially (see (Goodman, 2005, Section 6.6)). Hence, for all propagation distances larger than $\lambda$, diffraction in a homogeneous medium acts as a low-pass filter. Consequently, optical features of sub-wavelength resolution appear highly blurred in a microscope, due to the loss of information carried by their high spatial-frequencies. Over the years, numerous “hardware” methods for sub-wavelength imaging have been demonstrated (Lewis et al., 1984; Betzig et al., 1991; Di Francia, 1952; Lezec et al., 2002; Huang & Zheludev, 2009; Yildiz et al., 2003; Hell et al., 2009); however, all of them rely on multiple exposures. Apart from hardware solutions, several algorithmic approaches for sub-wavelength imaging have been suggested (see, e.g. (Harris, 1964; Papoulis, 1975; Gerchberg, 1974)). Basically, algorithmic sub-wavelength imaging aims to reconstruct the extended spatial frequency range (amplitudes and phases) of the information (“signal”), from measurements which are fundamentally limited to the range $[-n/\lambda, n/\lambda]$ in the plane-wave spectrum. However, as summarized by Goodman in his book (2005), “all methods for extrapolating bandwidth beyond the diffraction limit are known to be extremely sensitive to both noise in the measured data and the accuracy of the assumed a priori knowledge”, such that “it is generally agreed that the Rayleigh diffraction limit represents a practical frontier that cannot be overcome with a conventional imaging system.”

In spite of this commonly held opinion that algorithmic methods for sub-wavelength imaging are impractical (Goodman, 2005), a recent work proposed a method for reconstructing sub-wavelength features from the far-field (and/or blurred images) of sparse optical information (Gazit et al., 2009). The concept of sparsity-based
sub-wavelength imaging is related to Compressed Sensing (CS), which is a relatively new area in information processing (Candes et al., 2006; Candes & Tao, 2006; Donoho, 2006; Candes & Wakin, 2008; Duarte & Eldar, 2011). It has been shown that sparsity-based methods work for both coherent (Gazit et al., 2009; Szameit et al., 2010) and incoherent (Shechtman et al., 2010; Shechtman et al., 2011) light. An experimental proof-of-concept was presented in (Gazit et al., 2009): the recovery of fine features that were cut off by a spatial low-pass filter. Subsequently, these concepts were taken into the true sub-wavelength domain and demonstrated experimentally resolutions several times better than the diffraction limit: the recovery of 100 nm features illuminated by 532 nm wavelength light (Szameit et al., 2010). These ideas were followed by several groups, most notably the recent demonstration of sparsity-based super-resolution of biological specimens (Babacan et al., 2011).

Here, we take the sparsity-based concepts into a new domain, and present the first experimental demonstration of sub-wavelength CDI: single-shot recovery of sub-wavelength images from far-field intensity measurements. That is, we demonstrate sparsity-based sub-wavelength imaging combined with phase-retrieval at the sub-wavelength level. We recover the sub-wavelength features without measuring (or assuming) any phase information whatsoever; the only measured data at our disposal is the intensity of the diffraction pattern (Fourier power spectrum) and the support structure of the blurred image. Our processing scheme combines bandwidth extrapolation and phase retrieval, considerably departing from classical CS. We therefore devise a new sparsity-based algorithmic technique which facilitates robust sub-wavelength CDI under typical experimental conditions.

9.2 Sparsity based super-resolution

In mathematical terms, the bandwidth extrapolation problem underlying sub-wavelength imaging corresponds to a non-invertible system of equations which has an infinite number of solutions, all producing the same (blurred) image carried by the propagating spatial frequencies. That is, after measuring the far field, one can add any information in the evanescent part of the spectrum while still being consistent with the measured image. Of course, only one choice corresponds to the correct sub-wavelength information that was cut off by the diffraction limit. The crucial task is therefore to extract the one correct solution out of the infinite number of possibilities for bandwidth extension. This is where sparsity comes into play. Sparsity presents us with prior information that can be exploited to resolve the ambiguity resulting from our partial measurements, and identify the correct bandwidth extrapolation which will yield the correct recovery of the sub-wavelength image.

Information is said to be sparse when most of its projections onto a complete set of base functions are zero (or negligibly small). For example, an optical image is sparse in the near-field when the number of non-zero pixels is small compared to the entire field of view. However, sparsity need not necessarily be in a near-field basis; rather, it can be in any mathematical basis. Many images are indeed sparse in an
appropriate basis. In fact, this is the logic behind many popular image compression techniques, such as JPEG. In the fields of signal processing and coding theory, it is known for some time that a sparse signal can be precisely reconstructed from a subset of measurements in the Fourier domain, even if the sampling is carried out entirely in the low-frequency range (Vetterli et al., 2002). This basic result was extended to the case of random sampling in the Fourier plane and initiated the area of CS (Candes et al., 2006). An essential result of CS is that, in the absence of noise, if the “signal” (information to be recovered) is sparse in a basis that is sufficiently uncorrelated with the measurement basis, then searching for the sparsest solution (that conforms to the measurements) yields the correct solution. In the presence of noise (that is not too severe), the error is bounded, and many existing CS algorithms can recover the signal in a robust fashion under the same assumptions.

The concept underlying sparsity-based super-resolution imaging and sparsity-based CDI relies on the advance knowledge that the optical object is sparse in a known basis. The concept yields a method for bandwidth extrapolation. Namely, sparsity makes it possible to identify the continuation of the truncated spatial spectrum that yields the correct image. As was shown in (Gazit et al., 2009), sparsity-based super-resolution imaging departs from standard CS, since the measurements are forced to be strictly in the low-pass regime, and therefore cannot be taken in a more stable fashion, as generally required by CS. Therefore, a specialized algorithm was developed, Non Local Hard Thresholding (NLHT), to reconstruct both amplitude and phase from low-frequency measurements (Gazit et al., 2009). However, NLHT, as well as other CS techniques necessitate the measurement of the phase in the spectral domain. In contrast, the current problem of sub-wavelength CDI combines phase-retrieval with sub-wavelength imaging, aiming to extrapolate the bandwidth from amplitude measurements only. Mathematically, this problem can be viewed in principle as a special case of quadratic CS, introduced in (Shechtman et al., 2011). However, the algorithm suggested there is designed for a more general problem resulting in high computational complexity. Here we devise a specified algorithm that directly treats the problem at hand.

9.3 Sparsity based sub-wavelength CDI

For the current case of sub-wavelength CDI, the phase information in the spectral domain is not available. Hence, fundamentally, sub-wavelength CDI involves both bandwidth extrapolation and phase retrieval. However, despite the missing phase that carries extremely important information, we show that sparsity-based ideas can still make it possible to identify the correct extrapolation. Namely, if we know that our signal is sufficiently sparse in an appropriate basis, then from all the possible solutions which could create the truncated spectrum the correct extrapolation is often the one yielding the maximally sparse signal. Moreover, even under real experimental conditions, i.e., in the presence of noise, searching for the sparsest solution that is consistent with the measured data often yields a reconstruction that is very close to
the ideal one.

Our algorithm iteratively reveals the support of the sought image by sequentially rejecting less likely areas (circles, in the experiments shown below). Thus, the sparsity of the reconstructed image increases with each iteration. This process continues as long as the reconstructed image yields a power-spectrum that remains in good agreement with the measurements. The process stops when the reconstructed power spectrum deviates from the measurements by some threshold value. However, it is important to emphasize that the exact threshold value and the degree of sparseness of the sought image need not be known a priori, as our method provides a natural termination criterion. Namely, the correct reconstruction is identified automatically. A detailed description of the reconstruction method, as well as comparison with other methods (that do not exploit sparsity), are provided in Section 9.6.

9.4 Finding suitable basis

As explained above, sparsity-based CDI relies on the advance knowledge that the object is sparse in a known basis. In some cases, the “optimal” basis—the basis in which the object is represented both well and sparsely—is known from physical arguments. For example, the features in Very Large Scale Integration (VLSI) chips are best described by pixels on a grid, because they obey certain design rules. In some cases, however, the prior knowledge about the optimal basis is more loose, namely, it may be known that the object is well and sparsely described in a basis that belongs to a certain family of bases. For example, one may know in advance that the object is sparse in the near field using a rectangular grid, yet the optimal grid spacing is not known a priori. We address this issue in Section 9.8, where we describe a sparsity-based method that uses the experimental data to algorithmically find the optimal grid size (optimal basis) for our sub-wavelength CDI technique. That section also shows that the choice of basis functions is not particularly significant in our procedure: we obtain very reasonable reconstruction with almost any choice of basis functions, as long as they conform to the optimal grid. Finally, we note that recent work has shown that it is often possible to find the basis from a set of low-resolution images, using “blind CS” (Gleichman & Eldar, 2010). Likewise, in situations where a sufficient number of images of a similar type is available at high resolution, one can reconstruct the optimal basis through dictionary learning algorithms (Aharon et al., 2006).

9.5 Experiments

We demonstrate sub-wavelength CDI technique experimentally on two-dimensional (2D) structures. The optical information is generated by passing a $\lambda = 532$ nm laser beam through an arrangement of nano-holes of diameter 100 nm each. The sample is made of a 100 nm thick layer of chromium on glass; this thickness is larger than the
skin depth at optical frequencies, such that the sample is opaque except for the holes. We use a custom microscope (numerical aperture \( NA = 1 \), magnification \( \times 26 \)) and a camera to obtain the blurred image. The optical Fourier transform of the optical information is obtained by translating the camera to the focal plane of the same microscope.

The optical information is generated by passing a collimated laser beam (\( \lambda = 532 \text{ nm} \)) through a mask, whose transmission function corresponds to the optical information superimposed on the laser beam. The mask is fabricated as follows: As substrate material we chose fused silica, because it is a high quality transparent material at optical frequencies, and because its processing technology is well developed. In order to create a mask containing the optical image, we deposit opaque material on the substrate and make several patterned holes in it, such that the holes pass the light while the opaque material blocks it. For this purpose, we sputter a chromium layer onto the surface of the substrate. Chromium is a metal, which absorbs light at optical frequencies. Nevertheless, the thickness of the chromium layer has to be larger than the skin depth at optical frequencies, to avoid undesired transmission through that layer. Thus we select a thickness of 100 nm as suitable compromise between high quality optical behavior and fabrication considerations. The structures in the chromium layer are nano-holes, drilled in the chromium by a beam of focused gallium ions from a liquid metal ion source (Krohn & Ringo, 1975; Prewett & Jefferies, 1980) (Zeiss Neon 60). With this technology, it is feasible to mill the desired structures into the chromium layer directly and efficiently, without any additional lithography process. Utilizing a convenient set of parameters, it is possible to imprint the designed structures into the metal layer, without significantly affecting the substrate material, and with high spatial accuracy. We fabricated two different samples yielding a two-dimensional sub-wavelength optical structure: (a) a Star of David (SOD) image, consisting of 30 holes, with 100 nm diameter each, spaced by 100 nm; and (b) a “random” image comprised of 12 circular holes of 100 nm diameter each, placed in a random order. The Scanning Electronic Microscope (SEM) images of the samples are shown in Figure 9.1. Note that the SEM images are not in proportion as, in reality, the holes are of the same size and their diameter is equal to the spacing between holes. Generally, we use this approach throughout the paper: all images are shown in some abstract units that are, however, proportional to the corresponding physical quantities. The correspondence can be established using the fact that all holes are of diameter of 100 nm.
Let us present the reconstruction results first, followed by a full description of our method in Section 9.6. We begin with an ordered structure: a Star of David, consisting
of 30 nanoholes. Figure 9.2a shows an SEM image of this sample. Figure 9.2b depicts the image seen in the microscope. As expected, the image is small and severely blurred. The spatial power spectrum (absolute value squared of the Fourier transform) of the image is shown in Figure 9.2c. This truncated power spectrum covers a larger area on the camera detector, therefore facilitating a much higher number of meaningful measurements (each pixel corresponds to one measurement). We emphasize that only intensity measurements are used, in both the (blurred) image plane and in the (truncated) Fourier plane (Figures 9.2b, 9.2c, respectively), without measuring (or assuming) the phase anywhere. The recovered image, using our sparsity-based algorithm, is shown in Figure 9.2d. Clearly, we recover the correct number of circles, their positions, their amplitudes, and the entire spectrum (amplitude and phase), including the large evanescent part of the spectrum. This demonstrates sub-wavelength Coherent Diffractive Imaging: image reconstruction combined with phase-retrieval at the sub-wavelength scale. Moreover, as explained in Section 9.6, the intensity of the blurred image (Figure 9.2b) is used only for rough estimation of the image support. Our reconstruction method yields better results than other phase-retrieval algorithms (see comparisons in Section 9.7), because it exploits the sparsity of the signal (the image to be recovered), as prior information. As mentioned earlier, the underlying logic is to minimize the number of degrees of freedom, while always conforming to the measured data, which in this case is the truncated power spectrum (intensity in Fourier space). In the example presented in Figure 9.2, we take the data from Figures 9.2b and 9.2c, search for the sparsest solution in the basis of circles of 100 nm diameter on a grid, and reconstruct a perfect Star of David, as shown in Figure 9.2d. The grid is rectangular with 100 nm spacing (Section 9.8 describes how this parameter is found automatically), while the exact position of the grid with respect to the reconstructed information is unimportant (see Section 9.6).
We emphasize that our reconstruction algorithm is able to reconstruct the phase in the spatial spectrum domain (the Fourier transform), from the intensity measurement in Fourier space and some rough estimation of the image support. In addition, we use the knowledge that the holes are illuminated by a plane wave, implying non-negativity of the image in real space. In this Star of David example, our algorithm reconstructs...
the phase in the spectral plane, as presented in Figure 1e. For comparison, Figure 1f shows the phase distribution in Fourier space, as obtained numerically from the ideal model of the subwavelength optical information (calculated from the SEM image of Figure 9.2a). The reconstruction in Figure 9.2 therefore constitutes the first demonstration of subwavelength CDI.

Interestingly, when comparing the Fourier transform of the sample with the measured spatial power spectrum, one finds that more than 90% of the power spectrum is truncated by the diffraction limit, acting as a low-pass filter (see Figure 9.3). That is, we use the remaining 10% of the power spectrum and the blurred image, to successfully reconstruct the sub-wavelength features with high accuracy. In other words, the prior knowledge of sparsity and the basis is overcoming the loss of information in 90% of the power spectrum. As demonstrated in Section 9.6, it is the sparsity prior that makes it happen: without assuming the sparsity prior the reconstruction suffers from large errors.

![Image](image_url)

**Figure 9.3:** SOD image: available power spectrum.

The Star of David exhibits certain symmetries which could in principle assist the phase retrieval, had these symmetries been known. However, symmetry was not used for reconstruction of sub-wavelength features of Figure 9.2. Nevertheless, it is illustrative to present another example with no spatial symmetry at all: an irregular arrangement of sub-wavelength holes on the assumed grid. Figure 9.4a shows the blurred image of an unknown number of sub-wavelength circles, distributed in a random manner. The respective Fourier power spectrum, as observed in the microscope, is shown in Figure 9.4b. This sample is clearly not symmetric in real space, hence it does not exhibit a real Fourier transform. Still, we are able to reconstruct the sub-wavelength information, as shown in Figure 9.4c, where all features of the original sample are retrieved, despite the inevitable noise in the experimental system. Figure 9.4c shows the SEM image of the sample, displaying
the random arrangement of 100 nm holes. The electromagnetic (EM) field passing through these nano-holes has roughly the same amplitude for all the holes. The reconstructed amplitudes at the hole sites are represented by the colors in Figure 9.4c, highlighting the fact that the reconstructed field has similar amplitude at all the holes. The reconstructed phase in the spectral plane is presented in Figure 3e, where the white circle marks the cutoff imposed by the diffraction limit. As shown there, our algorithm recovers the phase throughout the entire Fourier plane, including the region of evanescent waves far away from the cutoff frequency. For comparison, Figure 9.4f shows the phase distribution in Fourier space, as obtained numerically from the ideal model of the sub-wavelength optical information (calculated from the SEM image of Figure 9.4d). Clearly, the correspondence between the original spectral phase and the reconstructed one is excellent, including in the deep evanescent regions. Interestingly, Figure 9.4e also displays the correct reconstruction of the phase around the faint high-frequency circle (of radius approximately 4 times the diffraction limit) where the phase jumps by $\pi$. Physically, this “phase-jump circle” is located at the first zero of the Fourier transform of a circular aperture, which in Fourier space multiplies the phase distribution generated by the irregular positions of the holes. The excellent agreement between Figures 9.4e and 9.4f highlights the strength of the sparsity-based algorithmic technique.
Figure 9.4: Reconstruction results for the "random" image.
9.6 Our sparsity-based reconstruction method for CDI

Under the experimental conditions described in the previous section, our problem amounts to the reconstruction of a signal from the magnitude of its Fourier transform, assuming furthermore that this information is known only for a small interval of low frequencies as shown in Figure 9.5. The discussion below is general and applies to both examples given in the paper (and, of course, to a very large class of optical images). However, in order to make the explanation more succinct, we demonstrate most of the results on the “random” image, because it has no implicit symmetries. The SOD image exhibits very similar behavior and its main results will be presented below.

![Figure 9.5: Fourier domain magnitude of the “random” image: (a) the full spectrum (simulated, without noise) needed to reconstruct the image precisely (by a simple application of the inverse Fourier transform), (b) the low-frequency part (actual measurements, in the presence of experimental noise).](image)

Of course, when the majority of the frequencies are lost, precise reconstruction is not possible, unless we have, or may assume, some additional information about the sought signal. In fact, the problem is even more difficult, because the measurements contain non-negligible noise. In a manner similar to (Gazit et al., 2009), we assume that the EM field in the object domain \((u,v)\) can be represented precisely, or approximated adequately (hereinafter, this relation is denoted by \(\cong\)) by means of a known generating function \(g(u,v)\). That is

\[
E(u,v) \cong \sum_m \sum_n x_{mn} g(u - m \Delta_u, v - n \Delta_v),
\]

where \(x_{mn}\) are unknown signal coefficients in the basis defined by the shifted versions of \(g(u,v)\). Note that the set \(\{(m \Delta_u, n \Delta_v)\}\) defines a rectangular grid where the shifted versions of the generating function are located. Hence, for example, by choosing
$g(u,v)$ to be the Dirac delta function we can obtain the sampled version of the continuous EM field distribution, where $\Delta_u$ and $\Delta_v$ define the sampling interval. Another classical example: all bandwidth limited signals can be represented precisely in this form when $g$ is chosen to be the jinc(\(\rho\)) function. For more examples see (Eldar & Michaeli, 2009) and references therein. Of course, the generator must be chosen in a way that corresponds to the signal in question (although, in the most general case of 2D information, the generator could simply be rectangular pixels). In this section and in Section 9.7, where we compare our algorithm with other methods, we assume that the basis function is chosen in a way that allows a perfect reconstruction of the sought signal, namely $g$ represents a circle of a priori known diameter (100 nm). We assume also that $\Delta_u = \Delta_v = 100$ nm. That is, we assume that the sought signal is comprised of non-overlapping circles of known diameter. The grid \{(m\Delta_u, n\Delta_v)\} containing all possible locations (144) is shown in Figure 9.6. Note that the exact placement of the grid is unimportant as our measurements are insensitive to shifts. A more detailed explanation of this property is presented in Section 9.8, where we discuss the implications of the grid assumption along with the impact of the basis function on the reconstructed signal.

Before moving on, there are two points we would like to stress. First, the assumption of an underlying grid is natural in many situations arising in digital signal processing. A prominent example are digital images that are comprised of pixels located on a rectangular grid. Just like in digital images, the grid in our case defines the resolution of a digitized version of the sought signal (see Section 9.8 for details). Second, it is important to note that all our comparisons with other methods are done under exactly the same assumptions, including a grid, basis functions, etc. As is evident from the experiments presented in Section 9.7, our algorithm outperforms other methods.

![Figure 9.6: The full grid.](image)

We emphasize that even if the correct number of circles were known (12 circles, in this example) there would be \(\binom{144}{12}\) > 10\(^{17}\) possible variants to choose from for the signal support. To limit the search space, we use the blurred version of the signal as
shown in Figure 9.7.

![Figure 9.7](image)

Figure 9.7: Support restriction by the low-resolution image: (a) blurred image magnitude, (b) grid restricted by the blurred image.

However, even after this restriction, there still remain \(\binom{37}{12} > 1.85 \times 10^9\) variants. More importantly, even after this restriction, the image cannot be reconstructed precisely unless additional information is available (see Section 9.7). Below we present our method that provides excellent reconstruction results based on the knowledge that the total number of circles in the image is small, that is, the image is \textit{sparse} in the basis associated with the circles, as defined by Equation (9.1).

In our method, we reconstruct the support and the magnitude of the circles in the sought signal simultaneously. To this end, we seek the sparsest \(x\) (\(x\) being a column vector comprised of the image coefficients \(x_{mn}\) as defined by Equation (9.1)), that yields a good agreement with the measurements. Mathematically, we try to solve the following optimization problem

\[
\begin{aligned}
\text{min} & \quad \|x\|_0 \\
\text{subject to} & \quad \|LFCx - r\|_2^2 \leq \epsilon, \\
x & \geq 0
\end{aligned}
\] (9.2)

Here \(\|\cdot\|_0\) denotes the \(l_0\) norm: \(\|x\|_0 = \sum_i |x_i|^0\), that is, \(\|x\|_0\) equals to the number of elements of \(x\) that are not zero. The measured (noisy) magnitude in the Fourier domain is denoted by \(r\). Note that the operators and inequalities, like \(|\cdot|\), and \(\geq\) are applied element-wise. The matrix \(C\) represents all possible shifts of the generator function (a circle); hence, \(Cx\) is the actual image that we reconstruct; \(F\) stands for the Fourier transform operator; \(L\) represents the low-pass filter. That is, \(L\) is obtained from the identity matrix of appropriate size by removing most of its rows while keeping only those that correspond to the low frequencies of its operand, as shown in Figure 9.5. Physically, \(L\) is the low-pass filter associated with the cutoff spatial
frequency of the optical system, which, for microscopes with NA=1, corresponds to the diffraction limit. Note that, due to errors in the measurements, the discrepancy in the Fourier domain is allowed to be up to some small value $\epsilon > 0$. A short discussion about the precise value of $\epsilon$ and whether it must be known a priori will follow. Note also that the last requirement $x \geq 0$ is valid because the optical information is generated by illuminating the sample with a plane wave, that is, a plane of equal amplitude and phase. Hence, the phase is the same across the whole image. Therefore, without loss of generality, we may assume that the phase is zero everywhere, since the absolute phase is unimportant. We do not assume that all circles have the same magnitude, however—they can have any value.

To solve (9.2) we developed an iterative method whose basic iteration contains the following two steps:

**Step 1:** Solve the minimization problem:

$$
\min \| |LFCx| - r \|_2^2 \\
\text{subject to } x \geq 0 .
$$

(9.3)

(in practice, we use an unconstrained formulation that is solved by the L-BFGS method (Liu & Nocedal, 1989)).

**Step 2:** After a solution $x$ to Step 1 is found, set to zero the entry of $x$ with minimal value. Once set to zero the entry remains so forever.

In theory, the iterations should be repeated so long as the constraint $\| |LFCx| - r \|_2^2 \leq \epsilon$ is satisfied. It is often argued that the value of $\epsilon$ is known a priori or can be estimated from physical constraints (as a matter of fact, in the case of the “random” image, the difference between the measured Fourier magnitude $r$ and its ideal variant $r^*$ is $\| r - r^* \|_2 = 1.7434$, which corresponds to a signal-to-noise ratio of $\| r^* \| / \| r - r^* \| = 1/0.041$). However, it is an important question whether the best value of $\| x \|_0$ (the true number of circles in the image) can be determined automatically. Consider the different stages of our method as shown in Figure 9.8. Is there any way to recognize that the correct number of circles is 12 without knowing $\epsilon$? It turns out that the answer to the above question is affirmative. As is evident from Figure 9.9, there is a big jump in the objective function value when the number of circles dips below the correct value of 12. Hence, even without knowing the noise bound $\epsilon$ one can easily identify that the smallest number of circles that “explains” well the measurements is 12 (this is, of course, correct as long as the circles have large enough amplitude). The result of our reconstruction and the true image are shown in Figure 9.10. Note that some circles have low magnitude so they are invisible in the color images. We therefore, place the ‘+’ sign at the center of all circles in the image ($x$’s entries that are not zeros).
Figure 9.8: Reconstruction stages for the example of the “random” image. Each stage (iteration) corresponds to a certain number of circles (non-zero entries in $x$):
(a) 37 circles, (b) 36, (c) 35, (d) 13, (e) 12, (f) 11, (g) 9, (h) 8, (i) 7.
Figure 9.9: “Random” image: objective function value (Fourier domain discrepancy) versus the number of circles in the solution.

Figure 9.10: Reconstruction result for the “random” image (a), and the true (original) image (b).

Very similar behavior is observed for the second image (SOD) whose results are shown in Figures 9.11 and 9.12.
Figure 9.11: SOD image: objective function value (Fourier domain discrepancy) versus the number of circles in the solution.

Figure 9.12: Reconstruction result for the SOD image (a), and the true (original) image (b).

In Section 9.8 we demonstrate that choosing an “incorrect” basis function, even one whose shape does not allow perfect representation of the sought signal, results,
nevertheless, in a reasonable reconstruction. Furthermore, we also demonstrate that the grid’s cell size can be determined automatically.

9.7 Comparison with other methods

We would like to stress again that our method is successful because we exploit the sparsity of the sought signal. To demonstrate this, we present a comparison with some classical reconstruction methods, and discuss the relation between our setup and classical compressed sensing.

9.7.1 Without a regularization

Our sparsity-based technique minimizes the $l^0$ norm subject to additional constraints. This formulation resembles closely a regularization imposed on $x$. Hence, the most naive approach would be to abandon the regularization altogether and to try to find $x$ that minimizes the discrepancy in the measurements. That is, we might solve the following problem

$$
\min \| |LFCx| - r \|^2 \\
\text{subject to } x \geq 0
$$

(9.4)

Note that this is exactly the problem we solve in the first iteration of our method. However, using this approach as the full reconstruction process has a number of drawbacks. First, the problem of image reconstruction from the magnitude of its Fourier transform (also called phase retrieval) is known to be particularly tough for continuous optimization techniques (for explanation and further details see (Osherovich et al., 2011)). To the best of our knowledge, the most widely used method for phase retrieval without additional information is the Hybrid Input-Output method (Fienup, 1982). A more detailed investigation of this method will follow in Section 9.7.3. Here, we present the results obtained by our optimization routine. As mentioned earlier, this formulation is equivalent to performing only one iteration of our method. Hence, the result is as shown in Figure 9.13. Note that the reconstruction contains many superfluous circles, and even if the correct number of the circles were known, a simple thresholding would yield an incorrect reconstruction.
Figure 9.13: Reconstruction without a regularization on $x$: (a) “random” image, (b) SOD image.

9.7.2 Replacing $l_0$ with another norm

Using $l_2$ regularization has long been a favorite among engineers due to its simplicity and the ability to obtain closed-form solutions in linear cases. In the non-linear case, these benefits are lost, of course. However, for us it is more important that the $l_2$ norm does not promote sparsity (actually, some papers claim that it usually results in the most dense solution possible (Chen et al., 1999)). To demonstrate that this regularization is not suitable for bandwidth extrapolation of sparse signals, we solved the following problem

$$\min \|x\| \quad \text{subject to} \quad \|LFCx - r\|^2 \leq \epsilon , \quad x \geq 0 . \quad (9.5)$$

The problem was solved by transforming it into an unconstrained optimization problem and choosing the weights of the penalty function terms so as to get the discrepancy in the measurements close to the true values. That is, assuming that the true $\epsilon$ is known ($\epsilon = 1.74$ in the case of “random” image, and $\epsilon = 0.0329$ in the case of SOD image). For the solution we used exactly the same routine (L-BFGS) as in our main algorithm. The results are shown in Figure 9.14.
Figure 9.14: Reconstruction using $l_2$ regularization: (a) “random” image, and (b) SOD image.

It is obvious that the reconstruction result is incorrect. Moreover, even if the correct number of circles were known, a simple thresholding would still produce an incorrect result.

Another viable alternative would be using the $l_1$ norm. A discussion on this norm is postponed to Section 9.7.4.

9.7.3 Methods based on alternating projections

In Chapter 3 we have already seen some methods for phase recovery (Gerchberg & Saxton, 1972; Fienup, 1982) that are based on a simple and elegant idea of alternating projections. Similar ideas was applied in the field of bandwidth extrapolation (Gerchberg, 1974; Papoulis, 1975). In general, the current signal estimate is transformed back and forth between the object and the Fourier domains. In each domain, all available information is used to form the next estimate. Here we consider two major methods of this type: Gerschberg type method (often referred to as Gerschberg-Saxton or Gerschberg-Papoulis) and Fienup’s Hybrid Input-Output method. As we already know, the former is a classical method of alternating projections where all available information in the current domain is imposed upon the current estimate. In the latter approach the object domain information is not directly imposed on the current estimate; instead a more complex update rule is used as we presented in Section 3.2.

Gerschberg type methods

As mentioned before, Gerschberg type methods are “pure” projection methods. The idea is to transform back and forth the current signal estimate between the signal and the Fourier domain performing a “projection” in each of the domains, that is, replacing the current estimate $x_{cur}$ with the nearest one that satisfies the constraints in the relevant domain ($x_{new}$). Hence, in each domain the following optimization
problem is solved

$$\min_{x_{new}} \| x_{cur} - x_{new} \|^2_2$$

subject to \( x_{new} \in S \), \hspace{1cm} (9.6)

where \( S \) denotes the set of all admissible signals in the current domain. In our case the current estimate is first Fourier transformed. Then the current (wrong) magnitude is replaced with the measured (correct) magnitude in the low-frequency regime. The resulting signal is back-transformed into the object domain (the result denoted by \( x' \)) where it is converted into an image comprised of circles (denoted by \( x_{new} \)) in the following manner. Recall that the image model is of the following form \( E = Cx \). Hence to find a projection we must solve the following problem

$$\min_{x_{new}} \| Cx_{new} - x' \|^2_2 ,$$

subject to \( x_{new} \geq 0 \). \hspace{1cm} (9.7)

The problem is convex and can be solved efficiently, however, we used a two-steps approximation instead of a full solution.

**Step 1** Solve \( \min_{x_{new}} \| Cx_{new} - x' \|^2_2 \). Note that this problem has a closed form solution: \( x_{new} = C^\dagger x' \), where \( C^\dagger \) denotes the Moore-Penrose pseudo-inverse of \( C \).

**Step 2** Set all entries of \( x_{new} \) that are negative to zero.

In general, this is not a true projection. However, it is a projection, if the vector \( x_{new} \) obtained after the first step is non-negative. This is indeed the case we observe in all our experiments. The results obtained after 5000 iterations of this method are shown in Figure 9.15. Usually, the correspondence between these and the true image falls considerably behind our sparsity-based reconstruction method.
From the results above, it is evident that the reconstruction is incorrect and even if the correct number of circles were known a simple thresholding would still result in incorrect images.

**Fienup’s Hybrid Input-Output method**

We already have met the Hybrid Input-Output method (see Section 3.2) that was developed by Fienup for the phase retrieval problem (Fienup, 1982). Although based on the method of alternating projections, HIO does not enforce the object domain constraints, that is, the image is allowed to be non-zero in the off-support areas and the values may be negative. To the best of our knowledge, HIO is the most successful numerical method for signal reconstruction from the magnitude of its Fourier transform. However, the method only achieves good results when all or most of the Fourier spectrum is available. Judging by the result shown below, the method is not suitable for the situation where the Fourier magnitude is available only for a small fraction of the frequencies. In our tests we applied the method in its original form, using only the Fourier domain magnitude and support information in the object domain (along with non-negativity). We did not try to enforce a constant value across every circle or zero values in the off-support areas, as the original method does not do that. As a post-processing step, the result returned by HIO was zeroed in the off-support areas (shown in Figures 9.16a and 9.17a) and then the values across each circle were averaged (shown in Figures 9.16b and 9.17b). As is evident from the results, the method is not capable of correct reconstruction of the signals. They cannot be recovered even if the correct number of circles is known: a simple thresholding will result in an incorrect reconstruction.
Figure 9.16: Fienup’s HIO method: “random” image results of reconstruction: (a) as produced by the method, (b) after enforcing a constant value across every circle.

Figure 9.17: Fienup’s HIO method: SOD image results of reconstruction: (a) as produced by the method, (b) after enforcing constant a value across every circle.

9.7.4 Relation to compressed sensing

Compressed sensing (CS) is an emerging field in image processing that performs signal reconstruction from a small number of its projections (Donoho, 2006; Candes et al., 2006; Candes & Tao, 2006). Conceptually, all CS techniques and their mathematical theory are based heavily upon the sparsity of the sought signals. It is important to note that CS, in its classical form, deals with measurements that are linear with respect to the unknown signal. Likewise, CS techniques generally assume random sampling distributed throughout the measurement domain. By contrast, in our current case of sub-wavelength CDI the measurements are: (1) nonlinear with respect to the sought
signal, (2) taken only in a small (low-frequency) region of the measurement domain, where (3) they are taken in a periodic fashion (dictated by the pixels’ arrangement of a digital camera sensor). Still, our reconstruction method relies on sparsity. As such, conceptually, our approach can still be viewed as CS in a broader sense.

Clearly, for the reasons stated above, many theoretical results and reconstruction methods of classical CS are not applicable to our problem. For example, the Matching Pursuit (MP) method (Mallat & Zhang, 1993) cannot be applied in its original form. Another popular method — Basis Pursuit (BP) (Chen et al., 1999), could, in principle, be applied here (considering BP as a general approach based on replacing the $l_0$ with the $l_1$ norm, rather than a specific algorithm). However, its benefits are not clear, because, in contrast to the linear case, in our nonlinear problem—using the $l_1$ norm still does not lead to a convex problem.

Besides the standard CS methods, which are inapplicable to the sub-wavelength CDI problem, it is instructive to consider other sparsity-based approaches which are related to CS, in the broader sense. One of these is based on division of the reconstruction process into two stages: at the first stage the missing Fourier phase is reconstructed using Fienup’s HIO algorithm (or Gerchberg-type method); at the second stage this phase is combined with the measured Fourier magnitude to form complete measurements that are linear with respect to the unknown signal. Once these linear measurements are available, one can use methods from classical CS (like, for example, BP) or our previously proposed method NLHT (Gazit et al., 2009), which is aimed at recovering data from low-pass measurements. We find, however, that this approach does not produce high quality results. This failure is, probably, attributed to inability of the projection-based methods to reconstruct the phase precisely, as shown in Figure 9.18 below.

![Figure 9.18: Fourier phase of the “random” image: (a) the true phase, (b) the phase obtained after 5000 iterations of HIO.](image)

Recently, several works have considered CS with quadratic nonlinear measurements (Shechtman et al., 2011; Candes et al., 2011). In both papers the resulting nonlin-
ear constraints are relaxed to semidefinite constraints using matrix lifting and an appropriate sparsity promoting objective is used. The work of (Candes et al., 2011) considers phase retrieval assuming the availability of several diffraction patterns obtained from multiple structured illuminations, which is not relevant to our problem. In contrast, the scenario considered in (Shechtman et al., 2011) is much closer to our current case. Namely, simultaneous phase retrieval and bandwidth extrapolation from a single-shot power-spectrum measurement. In fact, our present problem can be viewed as a special case of the problem addressed in (Shechtman et al., 2011). However, the algorithm suggested in (Shechtman et al., 2011) is targeting a more general problem, hence its computational complexity is high. With this reasoning in mind, we devised the new sparsity-based approach and algorithm described in the next section, which is tailored for the specific problem of sub-wavelength CDI.

9.8 A method for automatic grid determination, and the (un)importance of the basis function

In this section we would like to discuss the implications of our assumption regarding the existence of a grid that, in fact, defines a discrete set of allowed locations where the chosen basis function can be placed. In many cases, especially when the optical information represents experimental data, introducing such a grid is highly justified. For example, a digital image is obtained from a continuous intensity distribution by sampling it with a sensor that physically is an array of square pixels arranged in rows and columns. Hence, naturally, the grid is rectangular and the basis functions are squares whose size is equal to the grid’s cell size. Likewise, our reconstruction provides a digitized version of the true signal as if it were performed by a sensor whose pixels’ shape corresponds to the chosen basis function (circular in our experiments above). Hence, the grid used in our reconstruction algorithm essentially defines the resolution of the reconstructed image. This is especially true when the spatial extent of the basis function is smaller than (or equal to) the grid’s cell size. An example of such a sensor with circular pixels is shown in Figure 9.19.

However, there is an important dissimilarity between our case and the regular sampling in the object domain. Since our measurements contain only the Fourier magnitude and no information is available about the phase, we cannot distinguish between all the shifted versions of the original signal. That is, if $E(u, v)$ represents the original signal, our best hope is to reconstruct a shifted version of it, that is, $E(u - \Delta u, v - \Delta v)$ for some $\Delta u$ and $\Delta v$. Which version (shift) of the original signal is reconstructed depends, of course, on the reconstruction method. Because our method seeks the sparsest solution, we obtain the digitization that corresponds to the perfect alignment shown in Figure 9.19a and not the “misaligned” version shown in Figure 9.19b. Because in the latter case each circle in the original image “switches on” two pixels in the sensor, in contrast to one pixel per circle in the aligned case. Hence, one does not need to manually align the grid with respect to the sought signal.
as the best alignment is obtained automatically with our reconstruction method. The only concern regarding the grid alignment is related to the placement of the blurred image that we use for loose support estimation. Fortunately, the solution to this problem is easy: the blurred image must be placed in a way that guarantees maximal grid coverage, that is, we shall keep as many allowed locations as possible.
Figure 9.19: The sought signal (red) imposed on a sensor with circular pixels (blue). Note that the best alignment (a) is automatically obtained as it results in a sparser reconstruction than a bad alignment (b).

9.8.1 The impact of the basis function

Let us now consider the situations where the basis function is chosen in a way that does not allow a perfect reconstruction. Specifically, we consider basis functions in a
shape of a square and a triangle, as shown in Figure 9.20.

![Figure 9.20](image)

**Figure 9.20:** Basis functions that do not allow a perfect reconstruction: triangles (a), and squares (b).

As is evident from Figure 9.21, the reconstruction in these cases matches our expectations: we obtain the correct “digitized” version of the sought signal that corresponds to the chosen basis function and the grid. We emphasize the fact that all experiments are done with actual data that contains a significant amount of noise.
Figure 9.21: Reconstruction in the case of basis functions that do not match the sought signal.

Moreover, if we consider the progress of the reconstruction process (see Figure 9.22) we observe that even an incorrect choice of the basis function has no adverse effect on the reconstruction. This fact has a simple explanation: the difference between a circle and a square (or a triangle) of size 100 nm is much smaller than 100 nm. Hence, being able to distinguish between these shapes would mean effective resolution that is much better than 100 nm. Thus, we conclude that the shape of the basis function
is not of great importance so long as its size matches the size of a typical feature in the sought signal. In what follows, we evaluate the possibility to discover the most appropriate grid pitch (basis function size) automatically, without any prior information.

Figure 9.22: Reconstruction of the “random” image using different basis functions: objective function value (Fourier domain discrepancy) versus the number of circles/squares/triangles in the solution. Marker shape corresponds to the basis function shape.

9.8.2 A method for automatic determination of an optimal grid

So far, we have seen that the shape of the basis function has no severe impact on the reconstruction process. Moreover, the best possible alignment is obtained automatically due to our requirement of maximal sparsity. These two properties can be used for automatic determination of the optimal grid pitch. To this end we ran a series of experiments with different grids whose pitch varies from 10 to 32 pixels (corresponds to the range of 48–152 nm) using the square basis function of the size that matches the grid cell. As was mentioned earlier, the results of Section 9.8.1 show that the particular choice of the basis function is not very important. Hence, we could choose any shape of the size equal to the grid pitch. The choice of the square basis function was stipulated by the fact that most digital images are comprised
of square pixels. Hence, this basis function will, probably, be the first choice in the situation where nothing is known about the sought signal. For each grid pitch we ran a few iterations of our method keeping the lowest discrepancy in the Fourier space as a numerical value that corresponds to the current grid pitch. There is no need to solve the problem completely, as our goal here is to see whether the sought signal can be represented well by the current grid. We expect that fine grids (small pitch) will represent well the sought signal so long as the grid’s pitch is smaller than or equal to the size of a typical feature in the signal. However, once the grid becomes too coarse, we expect a rapid growth of the objective function value. Hence, we expect the graph to have the distinctive "L"-shape, similar to the graphs in Figures 9.22, 9.11, and 9.9. As is evident from Figure 9.23, our expectations are confirmed by the experimental results.

![Graph showing objective function value versus grid pitch size](image)

**Figure 9.23:** Objective function value (Fourier domain discrepancy) versus grid pitch size.

Note that the first sharp jump in the objective function value happens during the transition from 21 pixels (the correct value) to 22 pixels. However, it may be argued that the transition is not sufficiently apparent and the true value may lie in some small interval around 21 pixels. Hence, we evaluate the behavior of our reconstruction method for the grid pitch lying in the interval of 18–24 pixels. As is evident from Figure 9.24, only the correct value of 21 pixels results in a clear and
sharp jump after we dip below the correct value of squares (12). This property can be used for pinpointing the correct pitch size. Hence, an automatic subroutine for the optimal pitch determination is comprised of two steps: first, run a few iterations of our reconstruction method to obtain quantitative results indicating how well different grid sizes can represent the sought image; second, run a full reconstruction procedure for a limited range of pitches near the elbow in Figure 9.23 and check what pitch results in a clear evidence of existence of the sparsest solution (as in Figure 9.24).

Note that the obtained grid cell size is *optimal* in the sense that it satisfies two important properties simultaneously: first, it allows good approximation of the sought signal; second, it leads to a highly evident sparse solution.

The suggested method is also based on the sparsity assumption: it works well when there are a few features in the sought signal are of approximately the same size. This situation arises in many physical setups. However, we currently are working on extending the algorithm to cases where the signal features may be of various sizes.
Figure 9.24: Objective function behavior versus the grid pitch size (in pixels): (a) 21—the correct value, (b) 20, (d) 19, (f) 18, (c) 22, (e) 23, (g) 24.
9.9 Concluding remarks

In this chapter, we presented a technique facilitating reconstruction of sub-wavelength features, along with phase-retrieval at the sub-wavelength scale, at an unprecedented resolution for single-shot experiments. That is, we have taken coherent lensless imaging into the sub-wavelength scale, and demonstrated sub-wavelength CDI from intensity measurements only. The method relies on prior knowledge that the sample is sparse in a known basis (circles on a grid, in our examples). We emphasize that sparsity is what makes our phase retrieval work—the other assumptions used in the algorithm (non-negativity, bounded support and the known basis) alone are not sufficient. It is important to note that most natural and artificial objects are sparse, in some basis. The information does not necessarily have to be sparse in real space—it can be sparse in any mathematical basis whose relation to the measurement basis is known, for example, the wavelet basis or the gradient of the field intensity, given that this basis is sufficiently uncorrelated with the measurements. In all these cases our technique can provide a major improvement by “looking beyond the resolution limit” in a single-shot experiment. Since our approach is purely algorithmic, it can be applied to every optical microscope and imaging system as a simple computerized image processing tool, delivering results in real time with practically no additional hardware. The fact that our technique works in a single-shot holds the promise for ultrafast sub-wavelength imaging—one could capture a series of ultrafast blurred images, and then off-line processing will reveal their sub-wavelength features, which could vary from one frame to the next. Finally, we note that our technique is general, and can be extended also to other, non-optical, microscopes, such as atomic force microscope, scanning-tunnelling microscope, magnetic microscopes, and other imaging systems. We believe that the microscopy technique presented here holds the promise to revolutionize the world of microscopy with just minor adjustments to current technology—sparse sub-wavelength images could be recovered by making efficient use of their available degrees of freedom. Last but not least, we emphasize that our approach is more general than the particular subject of optical sub-wavelength imaging. It is in fact a universal scheme for recovering information beyond the cut-off of the response function of a general system, relying only on a priori knowledge that the information is sparse in a known basis. Our preliminary theoretical and experimental results indicate, unequivocally, that our method offers an improvement by orders of magnitude beyond the most sophisticated deconvolution methods. In a similar vein, we believe that our method can be applied for spectral analysis, offering a means to recover the fine details of atomic lines, as long as they are sparse (that is, do not form bands). In principle, the ideas described here can be generalized to any sensing/detection/data acquisition schemes, provided only that the information is sparse in a known basis, and that the measurements are taken in a basis sufficiently uncorrelated to it.
10 Afterword

An alternative title for this thesis could be “Prior information in the phase retrieval problem”. This also can describe the main line of the work presented here. In fact we discovered and showed how to use two powerful priors: approximately known Fourier phase, and sparsity of the sought signal. The material in each chapter represents an idea and the main results in a succinct form that is suitable for publication in a scientific journal. Therefore, some of the results were not included in this thesis. Part of them is available as technical reports listed below

- (Osherovich et al., 2010a)
- (Osherovich et al., 2010c)
- (Osherovich et al., 2009c)
- (Osherovich et al., 2008)

Other forms of prior knowledge, for example, defocused/blurred version of the sought signal can be found in (Osherovich et al., 2010b), which summarizes our work done in collaboration with KLA Tencor Inc.

We should also mention a standalone work done in (Osherovich et al., 2009a), where we used Fienup’s HIO algorithm to design an overcomplete dictionary in a way that its atoms (columns) are maximally uncorrelated. The method produces excellent results that are significantly better than the results produced by currently used algorithms.
Bibliography


Narkiss, G. and Zibulevsky, M. Sequential subspace optimization method for Large-Scale unconstrained problems. CCIT 559, Technion, EE Department, 2005.


Hessian (eigen decomposition) as a method of matrix decomposition. We focus on the bifurcation of a single moving wavefront, and we discuss how to use the hessian matrix to find the eigenvector. This development has been made in a recent paper, and it has led to the development of 5,000 eigenvalues and eigenvectors. The development of this technique is presented in Section 10, and it has been implemented in the computer science department. The development of this technique is presented in Section 10, and it has been implemented in the computer science department.

Technion - Computer Science Department - Ph.D. Thesis PHD-2012-04 - 2012
A summary of the text:

In the works of the author, new methods of numerical algorithms were developed for solving the well-known problem, in practice, this problem consists of reconstructing the missing value of the phase function. The problem of reconstruction is a problem that arises in many fields of physics, including, among others, astrophysics and crystallography. The technique of coherent diffraction imaging (CDI) and after that is created a new area in which it is possible to achieve very high resolutions, and the need for them increases with the development of new technologies.

In this area, the phase function is reconstructed from measurements taken using an objective lens and an electron microscope is used to detect a hologram. Under certain conditions, the hologram is equal to the phase function, but the pixel array of the CDI camera is sensitive only to the magnitude of the signal. For this reason, the problem is one of the problems where one must reconstruct a signal from a missing phase, which interferes with the objective lens.

In practice, one obtains a method that is not influenced by the type of microscope and an algorithm for numerical reconstruction. Consequently, the resolution is limited by the performance of the hologram. In other words, the theoretical resolution of the algorithm is limited by the performance of the hologram. This is different in the case of the hologram, since for a given hologram, the resolution is limited by the number of possible solutions. Depending on the conditions, there can be an infinite number of solutions, which can be used to obtain different representations of the hologram, or in cases where there are no holograms, or where the hologram is not available.

Alternating projections (alternating projections, alternating projections) are a mathematical tool for solving optimization problems, especially when it comes to complex variables. This approach is not specific to the problem of holography and can be used in various fields of physics and engineering. The approach is based on the idea of iteratively minimizing a function, which is a complex variable, and can be used to solve a wide range of problems.
המחקר נעשה בהנחיית פרופסור ירד בר וdoctor מיכאל ציבולבסקי ופובלקסי למודעי המструктор. אנימהו על טכניון עלה תמיכה כספית והדיבה בהשתלמותי. כמוכן, אנימהו על מערדה ימתה דרך פרופסור מ. שגב על תמיכתם כספית. ERC ולקרן לקלן (דרק פרופסור מ. שגב) על התמיכות כספית.
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חיבור על מחקר

לעט מיכלי חלקי של תרדימיות למחשב תיאור דוקטור בפילוסופיה

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