Measuring aberrations in the rat brain by coherence-gated wavefront sensing using a Linnik interferometer

Jinyu Wang, J.-F. Leger, Jonas Binding, A.C. Boccara, Sylvain Gigan, L. Bourdieu

To cite this version:

Jinyu Wang, J.-F. Leger, Jonas Binding, A.C. Boccara, Sylvain Gigan, et al.. Measuring aberrations in the rat brain by coherence-gated wavefront sensing using a Linnik interferometer. 2012. <hal-00716152>

HAL Id: hal-00716152
https://hal.archives-ouvertes.fr/hal-00716152
Submitted on 10 Jul 2012

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Measuring aberrations in the rat brain by coherence-gated wavefront sensing using a Linnik interferometer

Jinyu Wang1,2,3,4,5, Jean-François Léger1,2,3, Jonas Binding1,2,3,4,6, A. Claude Boccara4, Sylvain Gigan4,5* and Laurent Bourdieu1,2,3

1Ecole Normale Supérieure, Institut de Biologie de l’ENS, IBENS, Paris, F-75005 France
2Inserm, U1024, Paris F-75005 France.
3CNRS, UMR 8197, Paris, F-75005 France
4Institut Langevin, ESPCI ParisTech, CNRS UMR 7587, ESPCI, 1 rue Jussieu, 75005 Paris, France
5Fondation Pierre-Gilles de Gennes pour la Recherche, 29 rue d’Ulm, Paris, 75005 France
6Max Planck Institute for Medical Research, Jahnstraße 29, Heidelberg, 69120 Germany
*sylvain.gigan@espci.fr

Abstract: Aberrations limit the resolution, signal intensity and achievable imaging depth in microscopy. Coherence-gated wavefront sensing (CGWS) allows the fast measurement of aberrations in scattering samples and therefore the implementation of adaptive corrections. However, CGWS has been demonstrated so far only in weakly scattering samples. We designed a new CGWS scheme based on a Linnik interferometer and a SLED light source, which is able to compensate dispersion automatically and can be implemented on any microscope. In the highly scattering rat brain tissue, where multiply scattered photons falling within the temporal gate of the CGWS can no longer be neglected, we have measured known defocus and spherical aberrations up to a depth of 400µm.

©2011 Optical Society of America

OCIS codes: (110.1080) Active or adaptive optics; (010.7350) Wavefront sensing; (110.0113) Imaging through turbid media.

References and links

1. O. Albert, L. Sherman, G. Mourou, T. B. Norris, and G. Vdovin, "Smart microscope: an adaptive optics learning system for aberration correction in multiphoton confocal microscopy," Optics letters 25, 52-54 (2000).
2. P. Marsh, D. Burns, and J. Girkin, "Practical implementation of adaptive optics in multiphoton microscopy," Opt Express 11, 1123-1130 (2003).
3. M. J. Booth, M. A. Neil, and T. Wilson, "New modal wave-front sensor: application to adaptive confocal fluorescence microscopy and two-photon excitation fluorescence microscopy," J Opt Soc Am A 19, 2112-2120 (2002).
4. M. J. Booth, M. A. Neil, R. Juskaitis, and T. Wilson, "Adaptive aberration correction in a confocal microscope," Proc Natl Acad Sci U S A 99, 5788-5792 (2002).
5. D. Debarre, E. J. Botcherby, M. J. Booth, and T. Wilson, "Adaptive optics for structured illumination microscopy," Opt Express 16, 9290-9305 (2008).
6. D. Debarre, E. J. Botcherby, T. Watanabe, S. Srinivas, M. J. Booth, and T. Wilson, "Image-based adaptive optics for two-photon microscopy," Opt Lett 34, 2495-2497 (2009).
7. N. Ji, D. E. Milkie, and E. Betzig, "Adaptive optics via pupil segmentation for high-resolution imaging in biological tissues," Nat Methods 7, 141-147 (2010).
8. N. Ji, T. R. Sato, and E. Betzig, "Characterization and adaptive optical correction of aberrations during in vivo imaging in the mouse cortex," Proc Natl Acad Sci U S A 109, 22-27 (2012).
9. B. M. Hanser, M. G. Gustafsson, D. A. Agard, and J. W. Sedat, "Phase-retrieved pupil functions in wide-field fluorescence microscopy," Journal of microscopy 216, 32-48 (2004).
10. M. Rueckel, and W. Denk, "Properties of coherence-gated wavefront sensing," J Opt Soc Am A 24, 3517-3529 (2007).
11. M. Rueckel, J. A. Mack-Bucher, and W. Denk, "Adaptive wavefront correction in two-photon microscopy using coherence-gated wavefront sensing," Proc Natl Acad Sci U S A 103, 17137-17142 (2006).
12. M. Feierabend, M. Rueckel, and W. Denk, "Coherence-gated wave-front sensing in strongly scattering samples," Opt Lett 29, 2255-2257 (2004).
13. H. Schreiber, and J. H. Brüning, "Phase shifting interferometry," in Optical shop testing, D. Malacara, ed. (Wiley-Interscience, Hoboken, N.J., 2007), pp. 547-667.
14. S. Tuohey, and A. G. Podoleanu, "Depth-resolved wavefront aberrations using a coherence-gated Shack-Hartmann wavefront sensor," Opt. Express 18, 3458-3476 (2010).
15. J. Wang, J.-F. Leger, J. Binding, C. Boccara, S. Gigan, and L. Bourdieu, "Measuring aberrations in the rat brain by a new coherence-gated wavefront sensor using a Linnik interferometer," Proc. SPIE 8227, 822702 (2012).
16. J. Wang, J.-F. Léger, J. Binding, C. Boccara, S. Gigan, and L. Bourdieu, "Measuring known aberrations in rat brain slices with Coherence-Gated Wavefront Sensing based on a Linnik interferometer," in Biomedical Optics, OSA Technical Digest, p. BTu3A.83 (2012).
17. J. Binding, J. Ben Arous, J. F. Leger, S. Gigan, C. Boccara, and L. Bourdieu, "Brain refractive index measured in vivo with high-NA defocus-corrected full-field OCT and consequences for two-photon microscopy," Optics express 19, 4833-4847 (2011).
18. R. Crane, "Interference phase measurement," Appl. Opt. 8, 538–542 (1969).
19. D. C. Ghiglia, G. A. Mastin, and L. A. Romero, "Cellular-automata method for phase unwrapping," J. Opt. Soc. Am. A 4, 267-280 (1987).
20. R. Gens, "Two-dimensional phase unwrapping for radar interferometry: developments and new challenges," Int. J. Remote Sens 24, 703-710 (2003).
21. R. J. Noll, "Zernike polynomials and atmospheric turbulence," J. Opt. Soc. Am. 66, 207-211 (1976).
22. A. V. Larichev, P. V. Ivanov, I. G. Iroshnikov, and V. I. Shmal'gauzen, "Measurement of eye aberrations in a speckle field," Quantum Electronics 31, 1108-1112 (2001).
23. A. V. Koryabin, V. I. Polezhaev, and V. I. Shmal'gauzen, "Measurement of the thermo-optic aberrations of active elements based on yttrium aluminate and garnet," Quantum Electronics 23, 899-901 (1993).
24. M. Rückel, "Adaptive wavefront correction in two-photon microscopy using Coherence-Gated Wavefront Sensing," (Ph.D Ruperto-Carola University of Heidelberg, 2006).
25. Y. Piederriere, J. Cariou, Y. Guern, B. Le Jeune, G. Le Brun, and J. Lotrian, "Scattering through fluids: speckle size measurement and Monte Carlo simulations close to and into the multiple scattering," Optics Express 12, 176-188 (2004).
26. T. R. Hillman, Y. Choi, N. Lue, Y. Sung, R. R. Dasari, W. Choi, and Z. Yaqoob, "A reflection-mode confocal imaging system with enhanced light delivery and turbulence," Proc. SPIE 8227, 82271T (2012).
27. M. Oheim, E. Beaurepaire, E. Chaigneau, J. Mertz, and S. Charpak, "Two-photon microscopy in brain tissue: parameters influencing the imaging depth," J Neurosci Methods 111, 29-37 (2001).
28. D. Kleinfeld, P. P. Mitra, F. Helmchen, and W. Denk, "Fluctuations and stimulus-induced changes in blood flow observed in individual capillaries in layers 2 through 4 of rat neocortex," Proc Natl Acad Sci U S A 95, 15741-15746 (1998).
29. F. Helmchen, and W. Denk, "Deep tissue two-photon microscopy," Nat Methods 2, 932-940 (2005).
30. C. Wang, L. Qiao, F. He, Y. Cheng, and Z. Xu, "Extension of imaging depth in two-photon fluorescence microscopy using a long-wavelength high-pulse-energy femtosecond laser source," Journal of microscopy 243, 179-183 (2011).
31. D. Kobat, N. G. Horton, and C. Xu, "In vivo two-photon microscopy to 1.6-mm depth in mouse cortex," Journal of biomedical optics 16, 106014 (2011).
32. G. J. Tearney, M. E. Brezinski, J. F. Southern, B. E. Bouna, M. R. Hee, and J. G. Fujimoto, "Determination of the refractive index of highly scattering human tissue by optical coherence tomography," Opt Lett 20, 2258 (1995).
33. R. Juškaitis, "Characterizing High Numerical Aperture Microscope Objective Lenses," in Optical Imaging and Microscopy, P. Török, and F.-J. Kao, eds. (Springer-Verlag, Berlin Heidelberg, 2007), pp. 21-45.
34. M. J. Booth, M. A. A. Neil, and T. Wilson, "Aberration correction for confocal imaging in refractive-index-mismatched media," Journal of Microscopy 192, 90-98 (1998).
35. M. Feierabend, Coherence-gated wave-front sensing in strongly scattering samples (Ph.D Ruperto-Carola University of Heidelberg, 2004).
36. E. J. Botcherby, R. Juskaite, M. J. Booth, and T. Wilson, "An optical technique for remote focusing in microscopy," Optics Communications 281, 880-887 (2008).

List of abbreviations
NFP Nominal Focus Position; AFP Actual Focus Position; AO Adaptive Optics; BFP Back Focal Plane; CG Coherence Gate; CV Coherence Volume; dCV diffuse Coherence
Volume; DFT Digital Fourier Transform; CGWS Coherence Gated Wavefront Sensing; f focal length; FOV Field Of View; NA Numerical Aperture; eNA effective Numerical Aperture; OCT Optical Coherence Tomography; ff-OCT full-field Optical Coherence Tomography; OPD Optical Path Difference; PSI Phase Shifting Interferometry; SLED Superluminescent Light Emitting Diode; TPFM: Two-Photon Fluorescence Microscopy; SHS Shack-Hartmann Sensor; vSHS virtual Shack-Hartmann Sensor; MFP Mean Free Path.

1. Introduction

For biological imaging applications, light microscopy always compromises between image quality, penetration and non-invasiveness. However, due to refractive index mismatch (e.g. between water and tissue for water immersion objectives) and inhomogeneity within the specimen, wavefront distortions limit signal and contrast in deep imaging, especially for confocal, multi-photon microscopy or full-field Optical Coherence Tomography (ff-OCT). With an Adaptive Optics (AO) system, the wavefront of illumination can be shaped to compensate the distortion so that a diffraction-limited focus is restored. However, a traditional Shack-Hartmann sensor (SHS) can’t be used easily in strongly scattering biological samples since there is no selection of the ballistic photons originating from the reference point at a given depth in the sample amongst all scattered photons coming from the bulk.

To avoid this pitfall, several sensorless schemes have been implemented: (i) stochastic methods based upon genetic [1] or hill-climbing algorithms [2], in which strongly stained specimens and numerous iterations are needed; (ii) model-based approaches based upon modal wavefront sensing [3], in which the reduced number of measurements minimizes photobleaching and damage to the sample and which have been demonstrated in e.g. confocal microscopy [4], structured illumination microscopy [5] and two-photon fluorescence microscopy (TPFM) [6]; (iii) pupil segmentation methods [7, 8]; (iv) indirect aberration measurement through phase retrieval from images [9].

In contrast, coherence-gated wavefront sensing (CGWS) [10-12], which combined low-coherence phase shifting interferometry (PSI) [13] and a real SHS or a virtual SHS (vSHS) [10-12], can measure the distortions of the wavefront in scattering samples and therefore allow their correction by adaptive optics. Aberrations can be corrected at a high temporal rate since the wavefront distortions can be obtained with a single measurement. These pioneering studies implemented CGWS in TPFM, where a femtosecond pulsed laser was used as the low coherence light source and an appropriate amount of BK7 glass was incorporated in the reference arm for the compensation of dispersion between the two arms. CGWS has been validated in a solution of scattering beads [10, 12] and up to a depth of 200µm in the weakly scattering zebrafish forebrain [11]. Most recently, coherence gating with a real SHS has been demonstrated to reject unwanted back reflections [14] of lens surfaces far from the focus.

Here, we implement a new CGWS scheme based on a Linnik interferometer with a Superluminescent Light-Emitting Diode (SLED) as low-temporal-coherence light source. Compared to the previously described CGWS setup [10-12], its main advantages are the automatic compensation of dispersion between the two arms of the interferometer due to their symmetry and its possible implementation on any microscope due to its simple design [15, 16]. In fresh thick highly scattering rat brain slices, for a medium range numerical-aperture (NA) and low-magnification (20x/0.5) objective, we successfully measure up to a depth of about 400 µm a known defocus aberration, obtained by displacing the coherence gate (CG) position with respect to the actual focus position (AFP). With a high-NA and high-magnification (63x/0.9) objective, we measure up to a depth of about 200 µm the defocus as well as the 3\(^{rd}\) order spherical aberration introduced by the CG displacement. In contrast with the previous publications using CGWS [10-12], where backscattered light was mostly ballistic, multiple scattering is found to be not negligible in the rat brain. In addition to the ballistic photons, the CG of the SLED source also selects photons that have experienced
multiple scattering, if their time of flight lies within the temporal gate. The experimental results show that the multiply scattered photons selected by the CG begin to influence the wavefront measurements at shallow depths, of the order of the mean free path (MFP). However, we show that wavefront distortions can be measured at much larger depths, despite the fact that it relies on photons originating from a diffuse Coherent Volume (dCV), which is much larger than the Coherent Volume (CV) defined by ballistic photons only. CGWS measurement eventually fails, presumably when the dCV starts to deviate too much from the CV.

2. Materials and methods

2.1 CGWS setup

The experimental setup (Fig. 1) used to implement CGWS is a custom-made Linnik low-coherence interferometer (a Michelson interferometer with identical objectives in each arm) [15]. Two different pairs of water immersion objectives were investigated: medium range-NA and low-magnification (20x/0.5) objectives and high-NA and high-magnification (63x/0.9) objectives. The low-coherence source (SLED, center wavelength 750nm, bandwidth 23 nm) is collimated, made circularly polarized with the polarizer and the quarter wave plate, expanded to overfill the pupil of the objectives and split with a beamsplitter cube (BS1). The output of the interferometer is send to (a) CCD2 conjugated with the object plane by L1 (b) CCD1
conjugated with the pupil planes by the afocal telescope L1, L2 and (c) a real SHS (not shown on Fig. 1) also conjugated with the pupil planes. The SHS is used for calibration and alignment of each arm individually while the pupil of the other arm is blocked. CCD1 is used to record the interference pattern of the light fields on the pupils p1 and p2. Using PSI, the wavefront topography and hence the aberrations on the pupil p1 of the sample arm are measured. A grade A cover slip (refractive index 1.51) is placed on the rat brain slice to protect it. A similar cover slip is used in the reference arm to balance the dispersion and is slightly tilted to remove unwanted back reflections. A field diaphragm is placed between the two lenses of the afocal telescope to control and tune the microscope field of view (FOV). The FOV is calibrated for different values of the FD diameter both by using geometrical optics and by measuring it with an optical fiber translated in the focal plane of the objective.

The coherence length of our SLED is 10.8 µm (determined from its bandwidth), which corresponds to a CG length of 4.0 µm in the rat brain (refractive index 1.36 at 750nm estimated from [17]). The CGWS mostly senses the light scattered within the coherence volume (CV), whose axial length (centered at CG) depends on the coherence length, which is defined by the FWHM of the self-coherence function [10].

The sample can be shifted with TS1 along the optical axis, bringing different sample layers into focus. At the surface of the sample, the focus position coincides with CG position, by alignment (Fig. 2a). When imaging into a sample whose refractive index $n_s$ is greater than the index $n_i$ of the immersion medium, refraction at the surface causes the AFP of the objective to be shifted deeper into the sample with respect to the nominal focus position (NFP) (Fig. 2b). Because of the index mismatch between the refractive indices of water (used for immersion in the sample arm) and of the sample, the CV simultaneously moves in the opposite direction (Fig. 2b) [17]. By moving TS2 (Fig. 1), we can change the CV position and therefore either compensate this effect, or create a known additional defocus as well as some spherical aberrations (Fig. 2c) (see Appendix A).

In our coordinate system, the AFP is taken as the origin and the positive direction points deeper into the sample. We define the CG position as the distance between the AFP to the cross point of the CV center with the optical axis. Finally, as paraxial calculation is not valid for high-NA objectives, we locate the AFP with ray tracing method (see appendix A): a quasi point source of light is shifted along the optical axis and the AFP is defined as the position where the defocus aberration is zero (some higher order aberrations still remain).

Fig. 2. Principle of the method used to create known aberration. (a) Initial state at the surface of the sample (no aberration). (b) When focusing into the sample, the index mismatch moves the CV and the AFP in opposite directions away from the NFP and introduces tractable aberrations. (c) By changing the reference arm length, the CV position can be further displaced to add or subtract aberrations.
2.2 Wavefront reconstruction

In order to extract the phase of the electric field corresponding to the CGWS-selected backscattered light on CCD1 (Fig. 1), we use four-step PSI [13, 18] by shifting the mirror M mounted on the piezoelectric translation stage PZT (Fig. 1) in the reference arm. The mirror M is positioned at the focus of the objective during the alignment process using the SHS. The amplitude of the PZT is optimized to produce the optimal phase stepping of π/2. Theoretically (see appendix A), the PZT movements may introduce some defocus and high-order aberrations, but only up to a RMS wavefront error less than 1nm for both 20x/0.5 and 63x/0.9 objectives. We therefore omit this effect in our computation. The motion of the PZT is synchronized to the frame rate of CCD1. As the maximal frame rate of CCD1 is 60 frames per second (fps), the phase can be extracted at the maximum speed of 15 fps. For each measurement, the corresponding electric field (defined as the “CGWS signal”, and whose intensity is shown in Fig. 3b) is extracted from a recorded quadruplet of interferograms (Fig. 3a) by the four-step PSI algorithm.

Fig. 3. Wavefront reconstruction procedure. (a) Raw image recorded on CCD1 (imaging the objective pupils), in a rat brain slice, for AFP located 160µm deep below the coverslip and CG = -15µm from AFP (20x/0.5 objective). (b) Corresponding amplitude of the electric field obtained by PSI. Bottom right corner, schematic of the virtual lens array. (c) Intensity distributions in the virtual image plane of the sublens in white of subfigure (b) obtained by DFT for 4 CGWS images obtained at 4 neighboring sample positions. (d) The wavefront reconstructed from the slopes of the centroids of the vSHS at a given position and then averaged over 20 neighboring positions (here corresponding to a Zernike defocus coefficient of 0.31µm).
We use a vSHS [10, 12] to unwrap the phase extracted by PSI and reconstruct the actual wavefront [15]. Though the speckle arising in scattering samples prevents the use in CGWS of traditional phase unwrapping methods, which are strongly limited by path inconsistencies due to singularities [19] and discontinuities [20], the vSHS overcomes these problems and allows reliable wavefront unwrapping in the presence of speckle [10-12, 14]. The reconstruction of the wavefront was based on the algorithm implemented by Denk’s group [10, 12], in terms of the first 28 Zernike modes (up to 6th radial order) [21], which we briefly summarized here. The wavefront is numerically propagated through a virtual sublens array. For this purpose, the entire aperture is divided into a number of square-shaped sublenses (Fig. 3b). For the 20x/0.5 objective, we choose a sublens size of 20 by 20 pixels (49 sublenses across the pupil). For the 63x/0.9 objective, the sublens size was chosen as 13 by 13 pixels (47 sublenses across the pupil). For each sublens, we perform a discrete Fourier transformation (DFT) of the electric field over its sub-aperture. With scattering samples, the amplitude distribution of the electric field on a sublens is speckled. As a consequence, the DFT also shows a speckle structure in the focal plane [11, 22] (Fig. 3c). To reduce this speckle noise, we measure a number \( M \) (\( M=5 \)) of independent ensembles of scatters (Fig. 3c) [10, 11] corresponding to \( M \) positions of laterally shifted focus, placed on a grid with about 3\( \mu m \) spacing. At each position, (i) a center of mass algorithm [23] is used to locate the centroid of the intensity distribution, which is preferable to other techniques [22], (ii) the slopes of the local wavefront are calculated, (iii) the wavefront (described with Zernike coefficients) is reconstructed by least-square fit to the array of local slopes, (iv) the obtained individual Zernike coefficients are finally averaged over the \( M \) neighboring positions.

3. Results

3.1 Measuring known aberrations with CGWS in rat brain slices

In order to show that CGWS is able to measure aberrations in vivo, it is necessary to generate well-defined aberrations. The rat brain itself has some aberrations (that remain to be measured), but in order to assess the quality of our measurement, we have chosen to impose known aberrations and check that CGWS is able to measure them correctly. Previous work has shown that moving the position of the CG relatively to the AFP induces known aberrations [24]. This relative movement can either be produced by changing the length of the reference arm (fig. 2c) or simply by index mismatch when focusing in a tissue with refractive index different than water (fig. 2b). The two main aberrations introduced are defocus and 3rd order spherical aberration and their magnitude can be calculated exactly from the objective NA (Appendix A). Here, the following experiment is performed: (a) change the relative distance between the AFP and the CG for different depths in the rat brain; (b) calculate theoretically the expected aberrations; (c) measure the aberrations experimentally with CGWS and compare them to the theoretical predictions. Since the sample can itself create some unknown amount of aberration, we do not represent the absolute magnitude of the aberrations but their slope as a function of the relative position between the AFP and the CG.

We observe that CGWS accurately measures the defocus slope at shallow depths for both objective pairs, since the measured slopes match the theoretical predictions (Figs. 4a, 4b, 4d and 4e). When going deeper in the tissue, however, the measured slope is always smaller than the theoretical prediction, and rapidly diminishes (see e.g. Fig. 4d and 4e). The critical depth at which the drop occurs depends on the objective pair used. While for the 20x/0.5 objective we are able to measure the defocus down to 400 \( \mu m \) with little or no deviation, the accuracy degrades very much more quickly with the 63x/0.9 objective: at 300\( \mu m \), we measure already only half of the expected slope and the measurement tends to a flat curve with a zero slope.

We also compare the 3rd order spherical aberration measured by CGWS to predictions (Fig. 4c, 4f and 4g). For the 20x/0.5 objective, the expected spherical aberration slope is too small to be accurately measured by CGWS (it corresponds to a RMS wavefront error of \( \lambda/162 \))
when the CG is displaced by 10µm). However, for the 63x/0.9 objective, the theoretical 3rd order spherical aberration is much larger and CGWS is able to measure it accurately. Not only is the slope for shallow AFP depths close to the theoretical prediction, but also the drop in the measured slope occurs approximately for the same depth as for the defocus (compare Fig. 4e and 4g).

Finally, let us note that aberrations could not be measured at very shallow depths, for the AFP between 0 and 40µm. Even though our CV has a FWHM of only 10.8µm, its shoulders pick up the strong reflection from the glass surface, which dominates the backscattering from the tissue at the center of the CV.

Fig. 4. Measuring known aberrations at different depth for the 20x/0.5 and the 63x/0.9 objectives. (a) and (b): raw defocus measurement at different depths as a function of CG position for respectively the 20x/0.5 and 63x/0.9 objectives and theoretical curves. (c): raw 3rd order spherical aberration measurement for the 63x/0.9 objective at different depths as a function of CG position. In (a), (b) and (c), curves were vertically shifted for visibility. (d) and (e): from subfigures (a) and (b), defocus slope as a function of depth for respectively the 20x/0.5 objective and 63x/0.9 objective. (f) and (g): slope of the 3rd order spherical aberration and comparison with theory for respectively the 20x/0.5 objective and 63x/0.9 objective.

3.2 GCWS selects multiply scattered photons in rat brain slices

In order to understand what limits the maximum depth at which CGWS can be performed, we need a better understanding of the selection of photons by the CG as a function of the depth in the tissue. To get an insight into which photons actually fall within the CG and which information they carry about the wavefront of interest, we measure the speckle size and the magnitude of the CGWS signal at different depths for the two pairs of objectives (fig. 5).
The speckle size, which is calculated as the FWHM of the autocovariance function of the electric field on the pupil [25], is the most important parameter to assess the size of the coherence-gated region. Assuming ballistic propagation to and from the CV and a random distribution of scatters within the CV, we expect the speckle size to depend on the CV extension. If the size of the CV is of the order of the extent of the point spread function (PSF) of the objective and if the CV lies around the AFP, scatters within CV cannot be resolved by the objective and produce therefore a wavefront without speckle. If the CV moves away from the AFP, or if the CV extends laterally (as when aberrations are present or when multiple scattering starts to be non-negligible), then we expect to see in the CGWS signal a speckle structure, whose typical size is inversely proportional to the lateral extension of the CV. At the smallest depths, the size of the speckle is maximal when the defocus is minimal, as expected from the fact that the focus is the smallest at this position (Sup. Fig. 2).

Moreover, the speckle size also decreases monotonically with the depth in the tissue (Fig. 5a, FOV 1000µm and Fig. 5b, FOV 530µm). This can be interpreted as the fact that the effective coherent volume, from which selected photons seem to originate, is enlarged in depth by multiply scattering. We denote in the following this volume the “diffusive Coherence Volume” (dCV), which presumably is larger laterally than the CV and has an axial extension larger than the coherence length. Monte Carlo simulation have explored the spatial extent of this volume [26], but as exact scattering properties of the brain (in particular the anisotropy of the scattering) are still ill known, it is hard to predict the shape of this volume. The lateral extension of the dCV (d_{dCV}) (estimated from the typical size of the speckle in the objective pupil (d_{speckle}) by d_{dCV} ≈ 1.22 λ f_{obj}/d_{speckle}, where f_{obj} is the focal length of the objective) increases monotonically and seems to saturate at large depths (Fig. 5c, FOV 1000µm and Fig. 5d, FOV 530µm).

As the lateral extent of the dCV is of the order of several hundreds of µm at large depths, we checked whether the microscope FOV could limit the dCV and alter the CGWS measurement. When the CGWS system works, the FD (Fig. 1) is totally opened and it corresponds to the maximal microscope FOV (1000µm and 530µm for 20x/0.5 and for 63x/0.9 objectives respectively). By closing the FD, we measured the speckle size for several values of the FOV, calculated the corresponding dCV size (Fig. 5a, 5b, 5c and 5d) and computed the defocus slope to determine the FD range for which aberrations can still be estimated (Fig. 5e and 5f). As the FD is placed in the detection arm, it does not influence the geometry of photon trajectories entering in the sample. However, a photon exiting the sample with a large angle (as if it would come from a point outside the FOV) is blocked by the FD. Multiply scattered photons selected by the CGWS gate can therefore be rejected by the FD. Closing the FD reduces indeed the maximum dCV extension reached at large depth (Fig. 5c, 5d): some scattered photons in the dCV are blocked by the FD. This effect is present at large depths where multiply scattered photons contribute to CGWS signal. But perturbation of dCV by the FD is progressive and it can already be seen at depths smaller than the MFP, which is about 200µm in the cortex of adult rat in the far red [27-29]. The filtering of multiply scattered photons entering the CGWS gate by the FD is not precluding the correct estimation of the defocus slope (Fig. 5e and 5f), except for the very small range of FD size (FOV smaller than 180µm for the 20x/0.5 objective or than 60µm for the 63x/0.9 objective). Above these small FOV, the maximal depth at which defocus slope is correctly measured is the same for all FOV. Therefore, these curves show that the setup FOV does not determine the maximal depth of the CGWS measurement.

The effect of the depth on the CGWS signal is also an interesting parameter. In classical ff-OCT, the drop in signal is expected to follow the exponential attenuation of the ballistic light from Beer-Lamber’s law on a length scale given by half (because the light scattered at the focus travels twice the depth before exiting the sample) the MFP (~100µm in rat brain). In Fig. 5g and 5h, we represent the magnitude of CGWS signal as its total integrated intensity. The signal increases first until a depth of about 100 to 150 µm and then decays slowly with a
characteristic length scale larger than half the MFP (700µm for x20/0.5 objective, 250µm for the x63/0.9 objective). This dependence of the CGWS signal on depth could be accounted for by the selection within the coherence gate of a large fraction of snake-like multiply scattered photons. This confirms that the CGWS signal is influenced by multiply scattered photons in rat brain tissue.
Fig. 5. Transition from single scattering to multiple scattering in CGWS measurement within rat brain slices (male Wistar, 45 days old.) and influence of the microscope FOV. (a) and (b) Speckle size as a function of depth when CG is centered on AFP. (c) and (d) dCV lateral extension estimated from the speckle size as a function of depth when CG is centered on AFP. (e) and (f) Defocus slope as a function of depth. (g) and (h) The magnitude of CGWS signal as a function of the depth. (a), (c), (e) and (g) : 20x/0.5 objective; (b), (d), (f) and (h) 63x/0.9 objective. In each panel, the measurements are shown for four different diameters of the FD corresponding to four different FOV.

A possible explanation of the failure of CGWS at large depth could be that the speckle structure of the wavefront could not be resolved anymore at large depths by the camera pixels. To address this issue, we analyzed the wavefront at the maximal depths of measurement for each objective while binning the pixels of the camera (Fig. 6) before using the vSHS. We observe that for both objectives the defocus slope is still reliably measured, as long as there is at least about 1.2 to 1.3 pixels per speckle on average. It shows that the CGWS measurement is not limited by the camera sampling in our setup.

Discussion

Under the condition of single scattering regime, which is characterized by a ballistic transport to and from the CV and by a single backscattering event in CV, the wavefront distortions can be accurately measured by CGWS, as verified previously by [10-12]. We show in this paper, that for more strongly scattering structures such as the rat brain, multiple scattering starts playing a major role in CGWS. The photons selected by the temporal gate are not confined in the CV, but arise from a much larger volume referred to as the dCV.

The existence of an extended volume selected by temporal gating explains the CGWS performance as a function of depth. Due to multiple scattering, extra photons in the dCV participate in the wavefront measurement and the deeper in the sample, the larger the dCV. This effect (a) decreases the size of the speckle in the pupil (making the wavefront harder to measure) and most importantly (b) perturbs the wavefront itself, since it is averaged over photons coming from a much larger volume than CV.

Our main result is to demonstrate that the CGWS measurement of the known aberrations imposed in our experiment remains valid at large depths. CGWS could be performed indeed at depths much deeper than the depth at which only ballistic photons are selected by the CG. This result demonstrates the possibility to implement CGWS as wavefront sensor in a close loop system to improve microscopes working in scattering samples, such as two-photon microscopy or full-field OCT.
However, at depths larger than a critical depth, the CGWS fidelity drops. We have shown that this critical depth was neither due to a spatial filtering of the dCV due to a limited FOV of the microscope nor due to a poor sampling of the small speckle grain by the camera pixels. Therefore the maximal depth is more probably related to the fact that above the critical depth the wavefront averaged over all the photons in the dCV does not carry the same information as the one originating from the ballistic CV. A better understanding of this limit and of the fact that it depends on the objective used will need a model of scattering in the tissue [26].

It is interesting to compare CGWS with ff-OCT or conventional OCT. Our CGWS setup is mainly different in two aspects: the spatial coherence of the source plays a major role here and the coherence gating is performed directly in the pupil of high–NA objectives. In regular ff-OCT with thermal light, photons are indeed selected in the coherence window both based on their temporal and on their spatial position: the multiply scattered photons are mostly eliminated. In the CGWS setup, we measure the interference pattern with the reference in the pupil of the objective: we therefore select photons based only on their path length (the temporal coherence). As a consequence the multiply scattered photons that fall within this temporal gating participate in CGWS, even if they have been notably deviated. This results in a temporal gating only of the backscattered photons. We therefore select much more photons that just the ballistic ones.

Finally, in the regime determined above, where known aberrations could be reliably measured (200µm depth for the 63x/0.9 objective, 400µm depth for the 20x/0.5 objective), PSFs are calculated from the wavefront measured by CGWS using the first 28 Zernike coefficients whose tilt and defocus are numerically removed (Figure 7). It is calculated at the experimental CG positions where the defocus is minimal (but never exactly zero). We observe that the PSF degrades notably over the range of reliability of the CGWS. Some aberrations (mostly defocus and some amount of 3rd order spherical aberration) stem from the coherence gating itself, since the CG position was only close to the optimal defocus position. Other aberrations could have been introduced by the sample itself (e.g. 3rd order spherical aberration due to index mismatch [17]). However, these results have to be taken cautiously, in particular we cannot say anything about the isoplanatism of the aberrations as long as the wavefront we measured is not applied to correct a full-field image. However they give an estimate of the degradation of the image expected when going in depth in the sample.

Figure 7: Point Spread Function as extracted from the wavefront measured by CGWS for the minimal defocus position using the first 28 coefficients. The PSF was scaled over the whole gray values (0-255). We observe that the PSF degrades notably over the range of reliability of the CGWS.
5. Conclusion

We have implemented a new CGWS scheme based on a Linnik interferometer with a SLED as low temporal but high spatial coherence light source. Compared to the previous implementation of CGWS [10-12], its main technical advantages are the automatic compensation of dispersion between the two arms. Here the short coherence length for coherence gating is obtained from a broadband continuous source, with two symmetric arms to ensure that there is no phase delay between them for all wavelengths. Thanks to this simple design, a consequent advantage is its easy implementation on any microscope. Moreover, it offers in the future the possibility to modify easily the CGWS setup in terms e.g. of its central wavelength or its coherence length.

In fresh rat brain slices, we successfully measured up to a depth of about 400 µm for a 20x/0.5 objective and 200 µm for a 63x/0.9 objective known aberrations, obtained by displacing the CG position with respect to the AFP in the sample. However, measurement of the speckle size and the CGWS signal as a function of depth in the sample demonstrates that the CG was not successfully rejecting multiply scattered photons even at shallow depths. This was attributed to a large amount of multiply scattered photons, which could have similar time of flight in the sample as the photons of the reference arm. CGWS could be directly applicable at shallow depth or in thin slices, where regular wavefront sensing methods fail. It would allow the implementation of adaptive corrections. At larger depth, our results show that CGWS allows the quantitative measurement of known aberration despite the selection of a large amount of multiply scattered photons. Its benefits for close-loop adaptive optics have to be demonstrated by coupling it to any microscopy modality and by measuring the image improvement. Finally, imaging at large depth may require the improvement of the rejection of the multiply scattered photons e.g. by the increase of the scattering length e.g. by the use of higher-wavelength light [30, 31].

Acknowledgements

This work was supported by ANR RIB grants MICADO n° ANR-07-RIB-010-02 and ANR-07-RIB-010-04. J.B. and J.W were funded by the Foundation Pierre-Gilles de Gennes. We thank LLTech for the use of the LightCT software. We are grateful to Winfried Denk who allowed us to use the phase reconstruction algorithm implemented in his group.

Appendix A: Analytical computation under sine condition of the aberrations for a refractive-index-mismatch sample covered by a coverslip

For an objective operating under the sine-condition, an axial shift of the diffraction-limited focus corresponds to a defocus and to all orders of the spherical aberrations [32]. Binding et al. have proposed a concise formula to predict the aberrations for the mismatched index situation taking into account the spread between the NFP and AFP [17]. Here, we derive the analytical aberrations when a cover slip is inserted between the sample and the immersion water, as it is the case in most in vivo recordings when a glass coverslip is used to stabilize the brain after exposing it through a craniotomy.

Under the sine condition, the principal surface of the objective corresponds to a sphere segment of radius $n_i f$ (Fig.7) [33]. To determine the wavefront from a point A (i.e. the single pass aberrations) on the optical axis, we imagine a point source at this position and trace its rays into the back focal plane (BFP) of the objective. Ray tracing restricted to the meridional plane is sufficient because of rotational symmetry.

Emanating from A at an angle $\alpha_g$ with respect to OA, a ray is refracted and crosses the coverslip surfaces at P and B. The refraction angles in the glass coverslip and in the immersion water are $\alpha_g$, $\alpha_i$ respectively. It crosses the BFP at E (at a distance $r$ of the optical axis OZ). An auxiliary line BC is drawn perpendicular to the ray NC emanating from
the NFP (N) at an angle $\alpha_i$ with respect to OA. According to Fermat’s Principle, all rays of a planar wavefront transmitted through the objective have the same optical path length when they reach their common focusing point in the BFP. Thus all the rays at an angle $\alpha_i$ with respect to OZ in water arrive at point E, which is at a distance $r = n_i \sin \alpha_i f$ of OZ on the BFP.

For the light emanating from the NFP N in the aberration free case, a spherical wavefront will be produced at the principal surface, which corresponds to a planar one at the BFP. The optical path difference (OPD) $W(r)$ between the rays originating from A (in the presence of sample and coverslip) and from N (in the aberration-free case, i.e. in the absence of sample and coverslip, with the full light path in water) and reaching the BFP in E, is equal to the optical path from A to B minus the optical path from N to C.

Fig. 8 Geometry of schematic for ray tracing. Refractive index of water, glass slip, and sample are $n_i$, $n_g$, $n_s$, respectively. If the sample was a pure water solution and if there was no coverslip, the focus in this aberration free case would be located at the position N on the optical axis. The origin O is located at the cross point of the optical axis to the second surface of the glass coverslip, the Z axis is defined along the optical axis OA with the positive direction pointing towards the sample (away from objective). The distance ON is noted $d$, the thickness of the glass coverslip $T$, the actual point source is A and the distance AN is noted $\Delta Z$.

The optical path from A to B is:

$$OP(A) = n_s AP + n_g PB = n_s (d + \Delta z) / \cos \alpha_s + n_g T / \cos \alpha_g \quad (1)$$

The optical path from N to C is:

$$OP(N) = n_i NC = n_i [(d + T) / \cos \alpha_i + BD \sin \alpha_i]$$

where $BD = (d + \Delta Z) \tan \alpha_s + T \tan \alpha_g - (d + T) \tan \alpha_i$. Thus $W(r)$ will be:
\[ W(r) = (d + \Delta z)(n_s^2 - N_r^2)^{1/2} + T(n_n^2 - N_r^2)^{1/2} - (d + T)(n_i^2 - N_r^2)^{1/2} \] (3)

where \( N_r = \frac{r}{f} \) is the relative numerical aperture of the rays reaching \( E \) in the BFP.

Suppose the objective illuminated with a Gaussian intensity profile has an effective numerical aperture \( \leq NA \), see below) corresponding to a radius of the effective pupil \( r_0 \) in the BFP plane and an effective entrance angle \( \alpha_0 \), then the relative numerical aperture could be normalized as \( N_r = \frac{r}{eNA} \) or \( N_r = \frac{\sin \alpha_i}{\sin \alpha_0} eNA \). If \( T = 0 \), then \( W(r) = (d + \Delta z)(n_s^2 - N_r^2)^{1/2} - d(n_i^2 - N_r^2)^{1/2} \), which is same as [17]. If \( T = 0 \) and \( n_s = n_i \), \( W(r) = \Delta z(n_s^2 - N_r^2)^{1/2} \) which is same as [34-36].

Analytical Zernike coefficients can be computed by expanding \( W(r) \) into a series of Zernike modes by numerical convolution with the individual Zernike polynomials [21]. The AFP is defined as the position where the Zernike defocus is 0. For an objective that meets sine condition, all rotationally symmetric aberrations such as defocus, third order spherical aberration vary with CG position even when no actual aberrations are present [10]. When the CG position is moved with respect to the AFP, Eq. 3 is used to compute the Zernike defocus and the Zernike 3rd order spherical aberration as a function of CG position.

The aberrations created by moving the CG depends on the effective NA (eNA) of the objective illuminated with a Gaussian intensity profile. eNA was defined as the equivalent NA of an objective of same focal length \( f \) illuminated with a constant intensity profile and providing the same in-plane resolution. For an objective of pupil diameter \( D_p \), illuminated with a Gaussian beam of diameter \( D_b \) (measured at \( 1/e^2 \)), the truncation ratio \( t \) is defined as \( t = \frac{D_b}{D_p} \). Using [29], the 20X/0.5 objective (truncation ration 1.22) has an eNA of 0.47 and the NA 63X/0.9 objective (truncation ratio 1.44) has an eNA of 0.85.
Supplementary Information

1) CGWS setup

The components used are: a SLED light source (center wavelength 750nm, 1.2 mW, FWHM bandwidth 23 nm) coupled to a single mode fiber (NA = 0.11, EXS7505-841, Exalos, Swiss) with a collimator of focal length 10mm, AC080-010-B; Thorlabs, US); a polarizer Pol (NT47-603; Edmund Optics, US) and a 1/4 waveplate (AHWP05M-980; Thorlabs, US); beam expander (5x Zoom, BE02-05-B; Thorlabs, US); L1 (f = 150mm, G322352525, Linos, Germany), L2 (f = 200mm, G322353525, Linos, Germany); non-polarizing beam splitters BS1 and BS2 (50:50, BS014; Thorlabs, US); reference and sample arm pair of objectives (20x/0.5W, UMPlanFI; Olympus) or (63x/0.9W, HCX APO, Leica); a tilted coverslip identical to the one covering the sample to balance dispersion; a Piezo actuator PZT (resonant frequency 138kHz, range 9.1 ± 1.5µm, AE0505D08; Thorlabs, US) with a silicon mirror M (with low reflectivity in reference arm to improve the contrast of interference pattern) mounted on it; motorized linear translation stages TS1 and TS2 (range 28mm, T-LS28-M; Zaber, Canada); a field adjustable diaphragm; a CCD camera CCD2 conjugated to the objective focus (resolution 1024x1024, pixel size 14µm x14µm, pixel depth 12bits, DS11-01M15, Dalsa, Canada); a CCD camera CCD1 (resolution 1024x1024, pixel size 12µm x12µm, pixel depth 12bits, DS-21-01M60-12E, Dalsa, Canada). The pupil apertures of the sample and of the reference objectives are conjugated to the active surface of CCD1 by L1 and L2.

The beam dimensions are the following. The collimated beam of the SLED is circularly polarized after Pol and QWP. Before entering the interferometer, the beam is expanded to 11mm (1/e^2 width) by the beam expander, so that the objective pupils are overfilled by a factor of 1.22 for the 20x/0.5 objective and 1.92 for the 63X/0.9 objective. The diameter of the objective pupils imaged on CCD1 (12mm for the 20x/0.5 objective and 7.62mm for the 63x/ 0.9 objective) are smaller than the sensitive surface of CCD1 (12.28mm x 12.28mm).

2) CGWS Calibration

We calibrated the CGWS with a real SHS (HASO 3, Imagine Optic) using a mirror as sample and with the 20x/0.5 objective. By displacing the mirror along the microscope axis, we introduced a shift of twice the mirror displacement and the defocus aberration at each position was recorded both with SHS and CGWS. Both the SHS and CGWS correctly measured the slope of Zernike defocus and were in good agreement with the theoretical one (Supp. Fig. 1). The measured Zernike defocus coefficient of SHS has been normalized in the standard (Noll) notation [21].

Supp. Fig. 1. CGWS calibration. Measured defocus slope with SHS and CGWS compared to the predicted curve (for eNA=0.47 using the 20x/0.5 objective).
CGWS may report erroneous astigmatism (depending on the size of the scatters) if linearly polarized light is used [10], and these errors can be avoided by the use of circularly polarized light. Our setup, therefore, contains a quarter wave-plate in the sample arm (Fig. 1) to ensure that the sample was illuminated by circularly polarized light. We checked in this case that spurious astigmatism disappeared, thus demonstrating that no alignment-induced astigmatism was present in our system (data not shown).

3) Animal preparation

All surgical procedures were in accordance with the European Community guidelines on the care and use of animals (directive 86/609/CEE, CE official journal L358, 18th December 1986), French legislation (décret 97/748, 19th October 1987, J. O. République française, 20th October 1987), and the recommendations of the CNRS.

Before surgery, the rat was anesthetized by urethane injection (1.5 g/kg). Supplementary dose of urethane was applied if necessary. The brain was taken out, put into the solution (NaCL: 150mMol/L, KCl: 2.5mMol/L, Hepes: 10mMol/L pH = 7.4) and could be stored for 3 days at most in 4°C refrigerator. For CGWS measurement, the brain was sliced, then held on a glass slip, and covered with a cover slip to protect it. During the experiments, the slice was always immerged in the solution and the experiment lasted less than 3 hours.

4) Speckle dimension as a function of CG position at shallow depth.

At the smallest depths, the size of the speckle is maximal when the defocus is minimal, as expected from the fact that the focus is the smallest at this position (Sup. Fig. 2).

Supp. Fig.2. Speckle size as a function of the CG position relative to the AFP for different depths for the 20x/0.5 (a) and the 63x/0.9 (b) objectives.