Optics Letters

High resolution phase-sensitive sum frequency generation spectroscopy by timedomain ptychography

TOBIAS SCHWEIZER,² BRUNO G. NICOLAU,¹ PRISCILA CAVASSIN,¹ THOMAS FEURER,² NATALIE BANERJI,¹ AND JULIEN RÉHAULT^{1,*}

¹ Department of Chemistry and Biochemistry, University of Bern, Freiestrasse 3, 3012 Bern, Switzerland ² Institute of Applied Physics, University of Bern, Sidlerstrasse 5, 3012 Bern, Switzerland

*Corresponding author julien.rehault@dcb.unibe.ch

Received 22 July 2020; revised 11 September 2020; accepted 12 September 2020; posted 17 September 2020 (Doc. ID 403339); published XX Month XXXX

We demonstrate that time-domain ptychography, when applied to a set of broadband vibrational sum frequency spectra, reconstructs amplitude and phase of the vibrational free induction decay from an interfacial sample with a resolution that is independent of the up-converting pulse bandwidth and spectrometer resolution. These important improvements require no modifications to most standard homodyne setups and the method is applicable to other coherent homodyne spectroscopies like coherent anti-Stokes Raman spectroscopy or transient grating spectroscopy. © 2020 Optical Society of America

http://dx.doi.org/10.1364/OL.403339

Sum Frequency Generation Spectroscopy (SFG) is a method of choice to selectively investigate surfaces or interfaces [1]. In particular, vibrational SFG (vSFG) is used to study vibrational modes of interfacial molecules, providing structural and orientational information with sub-monolayer sensitivity. In the broadband version of vSFG (BBvSFG) a narrowband near infrared (NIR) pulse is mixed with a broadband infrared (IR) pulse that spans all vibrational frequencies of interest. The resulting sum-frequency signal from the interface is emitted in the phase matching direction (see Fig. 1.) and its spectral intensity I_{SFG} is typically measured with an array detector:

$$I_{SFG}(\omega_{SFG},\tau) \propto |E_{SFG}(\omega_{SFG},\tau)|^2 \tag{1}$$

where $\omega_{SFG} = \omega_{NIR} + \omega_{IR}$ is the sum frequency of the NIR and IR pulses at frequencies ω_{NIR} and ω_{IR} , respectively. The time delay τ is selected so that NIR probe and IR induced polarization overlap in time, but its exact value is somewhat arbitrary. Since the spectrum results from a homodyne measurement it contains only information on the vibrational amplitudes and phase information is lost. Yet, accessing the complex field $E_{SFG}(\omega_{SFG})$, specifically its phase, is crucial in order to determine correctly vibrational lineshapes or absolute orientations of interfacial molecules [2–8]. Hence, quite some effort has been dedicated to the development of Phase-Sensitive SFG (PS-SFG) modalities [8–16] which all rely on heterodyning the SFG signal with a local oscillator or the Non-Resonant Background (NRB) signal arising in some configurations[17,18]. These schemes successfully recover the complex field but come at the expense of a more complex experimental apparatus. In this letter, we demonstrate reconstruction of phase information from a series of homodyne measurements via time-domain ptychography [19], which requires no modification to a standard experimental setup. An algorithm is applied to a set of spectra measured at different time delays τ and reconstructs the complex linear vibrational polarization $P_S^{(1)}(t)$ from the interface. Moreover, we show that, in contrast to standard homodyne BB-vSFG measurements, the resolution of the reconstructed polarization is no longer limited by the spectrometer nor by the NIR bandwidth, but only by the largest delay τ . Thus, any standard homodyne BB-vSFG setup can produce phase information with unprecedented spectral resolution if a handful of spectra are measured at different delays τ and post-processed via time-domain ptychography. We named the new method HIPPY, for HIgh-resolution Phase-sensitive spectroscopy with PtYchography.



Fig. 1 (a) Experimental setup for BB-vSFG with an asymmetric probe pulse created via a Fabry-Pérot étalon. E_{NIR} , E_{IR} and E_{SFG} represent the fields of all pulse involved in the experiment, with their phase matching condition. (b) PIE scheme for the reconstruction of complex objects, here $P_{S}^{(1)}(t)$.

Time-domain ptychography is an emerging technique for phase and amplitude characterization of ultrashort coherent laser pulses [19]. In its basic implementation it uses the Ptychographic Iterative Engine (PIE) to reconstruct a complex object pulse from a set of sum-frequency spectra which result from the mixing of a well characterized probe pulse with an unknown complex object pulse at different delays between the two (see Fig. 1.)[19]. Briefly, the algorithm starts with a random complex object pulse as initial guess (see Fig. 1.) and in every iteration all measured spectra are processed. The algorithm calculates the product field for a particular time delay from the probe and the current estimate of the object pulse, calculates its Fourier transform and replace its modulus by the square root of the measured spectrum while preserving its phase. After an inverse Fourier transformation, the difference between the new and the old product field is used to update the current estimate of the object pulse. The best approximation to the actual object pulse appears typically after only a few iterations [20,21]. Time-domain ptychography has been successfully used to characterize attosecond or supercontinuum pulses [19,22-25], and the robustness of PIE has been demonstrated both theoretically and experimentally [26,27].

In a typical BB-vSFG setup a narrowband NIR pulse is mixed with a broadband IR pulse that covers all molecular vibrations of interest, typically between 800 and 4000 cm⁻¹. The center wavelength of the NIR pulse is selected such that the sum-frequency signal appears in a wavelength range that can be conveniently detected by silicon technology. Its bandwidth which determines the resolution of the recorded vibrational spectra, is typically adjusted via spectral filtering with bandpass interference filters, Fabry-Pérot étalons or zero-dispersion monochromators. In the impulsive limit of BB-vSFG, where the IR pulse is assumed to be short compared to the polarization, the signal field E_{SFG} is proportional to the second order polarization and can be approximated as [28,29]:

$$E_{SFG}(t,\tau) \propto P^{(2)}(t) \propto E_{NIR}(t-\tau) P_{S}^{(1)}(t)$$
 (2)

where $E_{NIR}(t - \tau)$ is the time delayed NIR pulse and $P_S^{(1)}(t)$ is the surface linear vibrational polarization or Free Induction Decay (FID) resulting from the interaction between the molecular sample through its complex linear susceptibility $\chi^{(1)}$ and the IR pulse $E_{IR}(t)$. Hereafter, we will use PIE notation and associate the $E_{NIR}(t - \tau)$ pulse with the probe and the polarization $P_S^{(1)}(t)$ with the complex

object to be reconstructed. To apply HIPPY to SFG spectroscopy we have to measure a set of homodyne SFG spectra for different time delays τ .

In our setup sketched in Fig. 1.(a) the NIR probe pulse is generated by passing a 100 fs pulse at 800 nm through a thin air spaced Fabry-Pérot étalon. The resulting narrowband probe is a time asymmetric pulse with a sharp rise and a slow exponential decay. Such probe pulses are conveniently used for mixing with long FIDs while removing the NRB that can arise from the overlap between NIR and IR pulses [30]. Here, the IR pulse was centered at 3.3 µm, corresponding to the energy of C-H stretch mode, and had a bandwidth of 240nm (200cm⁻¹). The probe pulse displayed in Fig. 2.(b) is calculated from the laser and Fabry-Pérot parameters, but also characterized with the integrated intensity of the SFG signal, which is dominated by the NRB and reflects the cross correlation between the NIR and IR pulses. The probe pulse can be delayed relative to the IR pulse via a mechanical translation stage. The sum-frequency spectra were recorded with a Kymera 193 spectrometer with a 1200 l/mm grating blazed at 600 nm combined with an Andor Idus 420 camera giving a spectral resolution better than 7 cm⁻¹ at 640 nm. For experimental demonstration, we used a monolayer of Octadecanethiol (ODT) on gold [31].

We measured seven spectra with a delay increment of 400fs and the spectral intensities between 2820 and 3020 cm⁻¹ are displayed in Fig. 2.(a) (see Data File 1, Data File 2, and Data File 3 [32]). The first two spectra exhibit a strong NRB contribution due to the gold substrate, which is nearly suppressed after delay #3. Also, the shape and intensity of the spectra and the vibrational signatures contained in them depend substantially on the time delay τ . While in standard BB-vSFG this constitutes a major obstacle for the correct interpretation of data, here all spectra are crucial as input for HIPPY.

The probe pulse and the seven spectra shown in Fig. 2(a) were subsequently used as input to the PIE algorithm (see Code 1 [33]). After every update the actual object, i.e. FID, was multiplied with the window function shown in Fig. 2(b) to remove noise in regions where no signal is expected. Typically, after only a few tens of iterations (about a second computation time) the algorithm had converged and reconstructed the spectra displayed in the lower panel of Fig. 2(a). Despite some minor discrepancies, the overall shape and magnitude of the spectra at different time delays is very well reproduced. The recovered vibrational FID is displayed in Fig. 2(b) and is Fourier transformed to



Fig.2. (a) Top: Intensity plot of measured spectra for different delays separated by 400 fs. Bottom: Corresponding reconstructed spectra. (b) Reconstructed FID (blue), probe pulse (red) and window function (dashed black curve) used for noise suppression. (c) Amplitude (blue) and phase (red) of reconstructed FID and the square root of the experimental spectrum for the first delay (dashed black curve). (d) Amplitude (blue) and phase (red) of $P_{S}^{(1)}$ after NRB subtraction and the square root of the experimental spectrum for the fourth delay (dashed black curve).

extract amplitude and phase of the frequency dependent signal. Note that absolute as well as linear phase (absolute time delay) cannot be recovered and would need additional reference measurements [34–36]. Therefore, we chose an arbitrary offset so that the phase is zero at the maximum amplitude and subtracted a linear phase so that the remaining phase is flattest for the spectral window considered. Here, only 7 spectra were sufficient for a high fidelity reconstruction, but we can easily include more spectra combined with a smaller time delay increment to improve the results, especially when the signal-to-noise ratio is lower. The spectral amplitude in Fig. 2(c) is dominated by a strong NRB, and the vibrational signatures from the ODT monolayer appear as dips in the NRB envelope.

Four important points are worth emphasizing. First, the retrieved signal does not depend on the shape of the NIR probe pulse. Second, the windowing effect in standard vSFG due to the finite duration of the NIR pulse is eliminated as the total time window is now determined by the number of recorded spectra times the time delay increment. Third, and this is a consequence of the previous points, the spectral resolution of the recovered signal is not limited anymore by the duration of the NIR pulse nor by the spectrometer resolution. To illustrate this, we compare the square root of the spectrum measured at the first time delay with the reconstructed amplitude in Fig. 2(c). Clearly, the reconstructed amplitude shows much better resolved vibrational modes and in fact now reveals their true lineshape. The improvement comes from the fact that time-domain ptychography merges the best of the frequency domain approach, i.e. acquiring highly resolved spectra at each acquisition with a sensitive CCD detector, together with the flexibility of the time domain approach, i.e. increasing the scan range by recording spectra at more and more time delays. Fourth, in BB-vSFG we typically must find a compromise between spectral resolution and signal strength. A longer probe permits a better resolution (if not limited by the spectrometer) but at the same time lowers the signal strength [3,8,37,38]. To circumvent this, alternative time domain techniques have been suggested. In our approach this is no longer an issue and we can easily use shorter probe pulses yielding higher signal strengths without sacrificing spectral resolution as long as we record enough spectra to cover the entire FID. Finally, we would like to comment on the NRB. For the data presented in Fig. 2(c) we chose to use all seven spectra, including the first two which feature a substantial NRB from the gold surface. Since HIPPY reconstructs whatever part of the object is scanned by the time-delayed probe pulses, we may in principle select only spectra taken at delays with a negligible NRB contribution (here after delay #4, see Fig. SI.1 for illustration). Alternatively, we can remove the NRB after the reconstruction. The latter is advantageous since the NRB heterodynes part of the signal and including the spectra with substantial NRB helps to improve the retrieval of the FID. To illustrate this, we subtracted the NRB from the FID, by fitting the NRB envelope and phase (assuming it to be flat in the middle of the spectrum of Fig. 2(c), see also Fig. SI.2). In Fig. 2(d) we show the resulting amplitude and phase of the FID that reflects now only the response from the ODT monolayer. The phase is displayed only were the amplitude is higher than 3% of the maximal amplitude. For comparison, we included the square root of the spectrum measured at the fourth delay, which is the first without NRB contribution. The comparison reveals three relevant points. First, the reconstructed vibrational frequencies for the three main peaks due to the terminal methyl group of ODT are more accurate and the modes exhibit narrower linewidths. Second, we identify spectral features that are not visible in the experimental spectrum, e.g. at 2845 cm⁻¹ and 2950 cm-1, which correspond to CH2 streching modes of ODT. Third, we find that the weak band around 2825 cm⁻¹ in the spectrum is absent in reconstructed FID, as it is an artifact due to probing with an asymmetric pulse and should be interpreted as such.



Fig. 3. Reconstructed linewidth as a function of a) total scan range, b) Fabry-Pérot linewidth and c) spectrometer resolution. Red arrows indicate parameters used for the simulation in Fig. 4.

To be able to reconstruct the complex polarization without being limited by the bandwidth of the probe nor by the resolution of the spectrometer is an important advantage that needs to be verified via numerical simulations. For these simulations we start with a single Lorentzian vibrational lineshape with a width of 1 cm⁻¹ sampled on a grid of 4'096 points with a spacing of 0.1 cm-1. The probe before passing through the air-spaced étalon is a 100 fs pulse at 800 nm and the delay increment is 150 fs. Fig. 3(a) shows the reconstructed linewidth versus the total time window scanned with an ideal spectrometer resolution and a Fabry-Pérot linewidth of 8 cm⁻¹. The longer the total scan window the better the match between reconstructed and original linewidth. Once the total time window is longer than the FID signal the reconstructed linewidth is constant and equal to the original linewidth. In Fig. 3(b) the Fabry-Pérot linewidth was scanned for a constant delay range of 100 ps and ideal spectrometer resolution. Up to approximately 25 cm⁻¹ the reconstructed linewidth matches the original. For larger values the Fabry-Pérot spacing becomes larger than the length of the laser pulse (100 fs) and the probe turns into a train of non-overlapping 100 fs pulses which causes the reconstruction to fail. Fig. 3(c) shows the reconstructed linewidth as a function of the spectrometer resolution for a constant delay range of 100 ps and a Fabry-Pérot linewidth of 8 cm⁻¹. Up to about 30 cm⁻¹ the reconstruction is perfect, above this value the spectral features blur out so much that the algorithm stops converging. Summarizing the results in Fig. 3 reveals that PIE reconstructs the true linewidth independent of étalon and spectrometer resolution up to values which are 30 to 40 larger than the original linewidth.

For the set of parameters indicated by red arrows in Fig. 3 (probe linewidth: 8.6 cm⁻¹, spectrometer resolution: 7 cm⁻¹ and scan range: 25 ps) we simulate the reconstruction of a more complex system containing four vibrational modes with different amplitudes, phases and linewidths. We used 30 spectra with a delay increment of 1ps. The comparison of the reconstructed amplitude and phase with the original indicates excellent agreement, as seen in Fig. 4. For example, the reconstruction of the 1.8 cm⁻¹ wide line results in a width of 1.9 cm⁻¹; the two lines separated by 3 cm⁻¹ are very well resolved (inset in Fig. 4) and the reconstructed phase is also very accurate over the entire spectrum. We would like to stress that all features mentioned above are narrower than the probe linewidth as well as spectrometer resolution as illustrated by the weakly resolved dashed line spectrum in Fig. 4.



Fig. 4. Reconstructed (blue) and input (red) amplitude (solid curve) and phase (dashed curve). The inset shows a zoom to the double peak at 2990 cm⁻¹ with a separation of 3 cm⁻¹. The black dashed line shows the square root of one of the simulated spectra.

Recovering the true linewidth only requires acquiring data at long delays with a sufficiently high signal-to-noise ratio.

In conclusion, we introduced HIPPY, a new method for PS-SFG spectroscopy that does not require heterodyning and has a spectral resolution which is independent of probe bandwidth and spectrometer resolution. Only a few spectra measured at different time delays between IR and NIR pulse in a BB-vSFG configuration are required. Processing these spectra with the PIE algorithm allows to reconstruct the complex FID from the interface with a spectral resolution that is limited only by the maximum time delay. A NRB contribution can help reconstruction, but it is not a prerequisite to the application of HIPPY.The PIE is fast, robust and easy to implement and can be applied to any coherent homodyne spectroscopy, in particular coherent anti-Stokes Raman spectroscopy and transient grating spectroscopy.

Funding. Schweizerischer Nationalfonds zur Förderung der Wissenschaftlichen Forschung (NCCR MUST); H2020 Mare Skłodowska-Curie Actions (801459, FP-RESOMUS); European Research Council (714586).

Disclosures. The authors declare no conflicts of interest

See Supplement 1 for supporting content.

REFERENCES

- 1. Y. R. Shen, Fundamentals of Sum-Frequency Spectroscopy (Cambridge Core 2016).
- 2. B. Busson and A. Tadjeddine, J. Phys. Chem. C 113, 21895 (2009).

3. I. V. Stiopkin, H. D. Jayathilake, C. Weeraman, and A. V. Benderskii, J. Chem. Phys. **132**, 234503 (2010).

4. A. D. Curtis, M. C. Asplund, and J. E. Patterson, J. Phys. Chem. C **115**, 19303 (2011).

5. Y. R. Shen, Annual Review of Physical Chemistry 64, 129 (2013).

A. L. Mifflin, L. Velarde, J. Ho, B. T. Psciuk, C. F. A. Negre, C. J. Ebben, M.
A. Upshur, Z. Lu, B. L. Strick, R. J. Thomson, V. S. Batista, H.-F. Wang, and F.
M. Geiger, J. Phys. Chem. A **119**, 1292 (2015).

7. W.-C. Yang and D. K. Hore, J. Chem. Phys. 149, 174703 (2018).

L. Fu, S.-L. Chen, and H.-F. Wang, J. Phys. Chem. B **120**, 1579 (2016).
N. Ji, V. Ostroverkhov, C.-Y. Chen, and Y.-R. Shen, J. Am. Chem. Soc. **129**, 10056 (2007).

10. N. Ji, V. Ostroverkhov, C. S. Tian, and Y. R. Shen, Phys. Rev. Lett. **100**, 096102 (2008).

11. S. Yamaguchi and T. Tahara, J. Chem. Phys. **129**, 101102 (2008).

12. J. Wang, P. J. Bisson, J. M. Marmolejos, and M. J. Shultz, J. Phys. Chem. Lett. **7**, 1945 (2016).

13. H. Vanselous and P. B. Petersen, J. Phys. Chem. C 120, 8175 (2016).

14. H. Wang, T. Gao, and W. Xiong, ACS Photonics 4, 1839 (2017).

15. M. Thämer, R. K. Campen, and M. Wolf, Phys. Chem. Chem. Phys. 20, 25875 (2018).

16. S. Sun, P. J. Bisson, M. Bonn, M. J. Shultz, and E. H. G. Backus, J. Phys. Chem. C 123, 7266 (2019).

17. O. Isaienko and E. Borguet, Langmuir 29, 7885 (2013).

18. O. Isaienko, S. Nihonyanagi, D. Sil, and E. Borguet, J. Phys. Chem. Lett. 4, 531 (2013).

19. D. Spangenberg, P. Neethling, E. Rohwer, M. H. Brügmann, and T. Feurer, Phys. Rev. A **91**, 021803 (2015).

20. B. C. McCallum and J. M. Rodenburg, Ultramicroscopy **45**, 371 (1992). 21. M. Guizar-Sicairos, K. Evans-Lutterodt, A. F. Isakovic, A. Stein, J. B.

Warren, A. R. Sandy, S. Narayanan, and J. R. Fienup, Opt. Express, OE 18, 18374 (2010).

22. D. Spangenberg, E. Rohwer, M. H. Brügmann, and T. Feurer, Opt. Lett., OL 40, 1002 (2015).

23. M. Lucchini, M. H. Brügmann, A. Ludwig, L. Gallmann, U. Keller, and T. Feurer, Opt. Express, OE **23**, 29502 (2015).

24. D.-M. Spangenberg, M. Brügmann, E. Rohwer, and T. Feurer, Appl. Opt., AO 55, 5008 (2016).

25. T. Witting, D. Greening, D. Walke, P. Matia-Hernando, T. Barillot, J. P. Marangos, and J. W. G. Tisch, Opt. Lett., OL **41**, 4218 (2016).

26. P. Sidorenko, O. Lahav, Z. Avnat, and O. Cohen, Optica, OPTICA **3**, 1320 (2016).

27. M. Murari, M. Murari, G. D. Lucarelli, G. D. Lucarelli, M. Lucchini, M. Lucchini, M. Nisoli, and M. Nisoli, Opt. Express, OE **28**, 10210 (2020).

28. J. E. Laaser, W. Xiong, and M. T. Zanni, J. Phys. Chem. B **115**, 2536 (2011).

L. Velarde and H.-F. Wang, Phys. Chem. Chem. Phys. **15**, 19970 (2013).
A. Lagutchev, S. A. Hambir, and D. D. Dlott, J. Phys. Chem. C **111**, 13645 (2007).

31. C. D. Bain, P. B. Davies, T. Hui. Ong, R. N. Ward, and M. A. Brown, Langmuir **7**, 1563 (1991).

32. T. Schweizer, B.G. Nicolau, P. Cavassin, T. Feurer, N. Banerji and J. Réhault, "Experimental SFG spectra"

https://boris.unibe.ch/id/eprint/146741, (2020).

33. T. Schweizer, B.G. Nicolau, P. Cavassin, T. Feurer, N. Banerji and J. Réhault, "Matlab Code for reconstruction of SFG spectra "

https://boris.unibe.ch/id/eprint/146748, (2020).

34. S. Sun, R. Liang, X. Xu, H. Zhu, Y. R. Shen, and C. Tian, J. Chem. Phys. 144, 244711 (2016).

35. P. E. Ohno, S. A. Saslow, H. Wang, F. M. Geiger, and K. B. Eisenthal, Nat Commun 7, 1 (2016).

36. J. M. Marmolejos, P. J. Bisson, and M. J. Shultz, J. Chem. Phys. **150**, 124705 (2019).

37. H.-F. Wang, Progress in Surface Science 91, 155 (2016).

38. J. P. Kraack and P. Hamm, Chem. Rev. 117, 10623 (2017).