Arbitrary-detuning asynchronous optical sampling pump-probe spectroscopy of bacterial reaction centers
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Asynchronous optical sampling (ASOPS) is a widespread technique for rapid time delay scanning, which has found numerous applications, e.g., in time-resolved pump-probe or Fourier-transform spectroscopy. This stroboscopic approach usually relies on two femtosecond oscillators of nearly identical [1–7]—or nearly multiple [2,8]—repetition rates. As compared to more conventional scanning methods based on a mechanical translation stage, the advantages of ASOPS include a faster scanning speed, the removal of pointing instabilities and defocusing for large time delays, the reduction of mechanical noise, and the possibility of achieving longer time measurement windows [1–7].

We have recently reported a new variant of ASOPS, coined arbitrary-detuning ASOPS (AD-ASOPS), which can be more easily applied to any pair of free-running femtosecond oscillators thanks to the lack of any specific requirement on the laser respective repetition rates [9]. In this Letter, we take advantage of this new feature to achieve asynchronous sampling combining for the first time, to the best of our knowledge, a chirped pulse oscillator as the pump laser, thus extending the available time window to almost 200 ns with a time resolution as good as about 320 fs. The method is illustrated with the measurement in a single experiment of the complete charge transfer dynamics of the reaction center from Rhodobacter sphaeroides. © 2013 Optical Society of America

Figure 1 shows the experimental setup. The pump laser is a CPO (Femtosource XL500, Femtolasers), delivering 4 nJ, 15 fs pulses at a repetition rate of 74.7 MHz, while the probe laser is a chirped-mirror Ti:sapphire oscillator (Synergy PRO, Femtolasers), delivering 530 nJ, 50 fs pulses at a repetition rate of 5.1 MHz. Since the two pump-probe delay is computed from the knowledge of the number of probe pulses that elapsed since the last pump-probe coincidence event, associated with the number of pump and probe pulses between the previous and next coincidence events. The data are then averaged in the corresponding time bin so that a complete time-resolved dynamics can be obtained after appropriate data averaging. In a previous AD-ASOPS experiment, we have demonstrated a time resolution of about 400 fs, limited by the uncertainty in coincidence detection and by the deviation from the assumed linear evolution resulting from laser timing jitter [9].

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Figure 1. Experimental setup for AD-ASOPS pump-probe spectroscopy, including the free-space pump-probe experiment and the fiber-based coincidence detection setup. The inset shows the setup used for characterizing the time resolution of the experiment, where a nonlinear BBO crystal is used in place of the sample.
lasers, both centered at 800 nm, have largely overlapping spectra, coincidence detection can be easily achieved using a fiber-based setup already described elsewhere [9]. The latter measurement relies on the linear interference between the two lasers and is thus highly sensitive, so that only a small fraction of the incident energy is required and is easily provided using the leaked beam transmitted through high-reflection dielectric mirrors. Two interference filters placed before the optical fiber couplers allow enhancement of the spectral overlap, resulting in a coherence time of 210 fs full width at half-maximum (FWHM), assuming a Gaussian profile as deduced from the measured spectra. The coincidence detection threshold, as described in [9], was fixed at 25% of the maximum interference signal, which results in a theoretical resolution limit of 295 fs. The cross-polarized pump and probe pulses are then focused on the sample, and the transmitted probe pulses are individually measured using a photodiode placed after an analyzer rejecting the scattered pump radiation. For each laser shot, the transmitted probe energy and the coincidence signal are acquired simultaneously using a 210 MS/s, 14-bit analog-to-digital acquisition card (X5-210M, Innovative), clocked at the probe-laser frequency.

Note that using a pump laser repetition rate as low as 5.1 MHz results in a significant decrease of the coincidence rate as compared to our previous work [9], increasing the time interval between coincidence events and thus increasing the detrimental effect of timing jitter on time resolution. In order to investigate this effect, we first characterized the time resolution by replacing the sample with a type-I BBO crystal, as shown in the inset of Fig. 1, allowing measurement of the cross-correlation signal and hence of the time resolution of the AD-ASOPS pump-probe setup.

Figure 2 shows the measured autocorrelation curves corresponding to data associated with coincidence pairs separated by less than 1 ms (red diamonds) or less than 0.5 ms (blue circles). Indeed, although the average time between coincidence pairs is roughly equal to 5 ms, the random nature of coincidence events resulting from the mechanical behavior of the two free-running oscillator cavities, associated to a roughly exponentially decaying distribution [9], allows selection of more favorable coincidence pairs. Selecting only coincidence pairs closer than 0.5 ms (blue dots) results in a time resolution of about 320 fs, as deduced from the FWHM of the Gaussian fit. This value is very close to the theoretical limit of 295 fs due to the uncertainty in coincidence detection, indicating that the drift in cavity length during a 0.5 ms time interval can be safely neglected. In contrast, the data associated with a time interval of 1 ms between coincidence pairs results in a time resolution greater than 600 fs, evidencing the detrimental effect of timing jitter. It is noteworthy that, due to the more restrictive conditions for the first measurement, the acquisition time is about three times longer for the first experiment than for the second one, so that one must compromise between time resolution and acquisition time (or signal-to-noise ratio).

We applied the AD-ASOPS pump-probe setup shown in Fig. 1 to investigate the dynamics of the \( P^+H_L^- \) charge separated state intermediate in bacterial photosynthesis, by monitoring the formation and decay of the electrochromic band shift of the B band, at 780 nm, close to its induced absorption maximum [12]. The experiment was performed in isolated carotenoid-containing wild-type \( R. \) sphaeroides RCs, in which the quinone acceptor \( Q_A \) was precluded. Under these conditions, the build-up of any long-lived (>200 ns) triplet states of the bacteriochlorophyll dimer electron donor P is avoided by efficient quenching by the carotenoid [13]. The 1 mm optical path length sample was placed in a sample holder that was thermostatted at 10° C and continuously rotated at ~3 Hz (diameter = 8 mm) in order to minimize sample degradation. The RCs were purified as described elsewhere [14], and were prepared to OD at 800 nm ~0.5 in 20 mM Tris, pH 8.0, 0.1% LDAO, using 10 mM dithiothreitol as a reductant. The energy of the 50 fs, 800 nm, s-polarized pump pulses was adjusted to 2 nJ, while the energy of the 15 fs, 800 nm, p-polarized probe pulses was adjusted to 50 pJ. The two beams were focused using an optical lens of focal length 100 mm, and the transmitted probe beam was spectrally filtered using a low-pass filter rejecting wavelengths longer than 800 nm in order to avoid averaging out of the signal around the isosbestic point. The resulting probe spectrum was centered at 780 nm with a spectral width of 10 nm FWHM.

Figure 3 shows the differential absorbance signal obtained using the experimental approach outlined above, using a total acquisition time of only 4 min. The data recorded in the time interval [~20 ps, 40 ps] were subject to a selection keeping only coincidence pairs separated by less than 1 ms (corresponding as discussed above to a time resolution of 600 fs), hence the greater amount of noise due to the smaller number of samples. The data are plotted using a linear-logarithmic scale extending up to 196 ns, exhibiting a large variety of time scales in this system. The solid line is a three-exponential fit that yields time constants equal to 2.0 ns (53%), 8 ns (42%), and 43 ns (5%). Our results are in good agreement with previously reported results obtained using low-repetition-rate continuous-pulse experiments [15,16], where the multiphasic kinetics were interpreted in terms of a gradual stabilization of the \( P^+H_L^- \) state. Our high time

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**Figure 2.** Cross-correlation signal obtained from sum-frequency mixing in a type-I BBO crystal. The blue circles (resp. red diamonds) are the data measured by selecting coincidence events closer than 0.5 ms (resp. 1 ms). The solid lines correspond to Gaussian fits, indicating a FWHM time resolution of 318 fs (resp. 606 fs).
resolution and excellent signal-to-noise ratio furthermore allow us to more precisely determine the rate and relative amplitude of the fastest component, which presumably reflects recombination from the unrelaxed state. Additionally, our experiment reveals a rise time with a time constant of 3.7 ps, also in agreement with previous measurements obtained using conventional ultrafast pump-probe spectroscopy with a mechanical delay line\( ^{12,17} \).

To summarize, we have applied AD-ASOPS to pump-probe spectroscopy in a biological system using a 74.7 MHz oscillator combined with a 5.1 MHz CPO. The relatively low repetition rate of the CPO made possible the application of ASOPS to a broad time window extending up to 200 ns, while keeping a best time resolution of about 300 fs. We were thus able to measure in a single experiment the complete charge separation and recombination dynamics of carotenoid-containing \textit{R. sphaeroides} RCs. Our results are in good agreement with previous experiments reported earlier but using different experimental approaches suited either to long \( ^{15,16} \) or short \( ^{12,17} \) time scales. Furthermore, our results exhibit an improved signal-to-noise ratio despite an acquisition time of only a few minutes. Finally, we stress that the large energy per pulse available in CPO lasers makes the generation of new spectral components straightforward, thus opening new perspectives for AD-ASOPS pump-probe spectroscopy in a great variety of spectral domains.

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### References and Notes

11. Since the time resolution of the experiment—limited by other factors discussed in the text—is more than five times longer than the specified laser pulse durations, an accurate measurement of the actual pulse durations at the time of the experiment was considered unnecessary.