Characterizing dissociative motion in time-resolved x-ray scattering from gas-phase diatomic molecules

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Time-resolved x-ray scattering (TRXS) measures internuclear separations in a molecule following laserinduced photoexcitation. The molecular dynamics induced by the excitation laser may lie on one or several bound or dissociative electronic states. Time-resolved x-ray scattering from these states can be difficult to isolate because they generally overlap in the angle-resolved x-ray scattering pattern $I(x, y, \tau)$, where τ is the pump-probe delay and (x, y) are the physical pixel positions. Here we show how standard transform methods can isolate the dynamics from individual states. We form the temporal Fourier transform $\tilde{I}(x, y, \omega) = \int_{-\infty}^{+\infty} d\tau \, e^{-i\omega\tau} I(x, y, \tau)$. This frequency-resolved x-ray scattering (FRXS) signal segregates the bound states according to their vibrational frequencies ω_i and also displays dissociative states along straight lines $\omega = vQ$, where the slope v is the rate of increase of the internuclear distance and Q is the momentum transfer between the incident and scattered x-ray photon. We derive this relation and use FRXS to extract state-specific dynamics from experimental TRXS from molecular iodine following a 520-nm pump. Dynamics observed include one- and two-photon dissociation of the ${}^{1}\Pi_{u}$ and ${}^{1}\Sigma_{e}{}^{+}$ excited states and vibrational wave packets on the $B{}^{3}\Pi_{0u}{}^{+}$ state.

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I. INTRODUCTION

The ability to observe time-resolved motion of electrons and nuclei in molecules is one of the principal goals of femtochemistry [1]. Time-resolved x-ray scattering (TRXS) and time-resolved electron scattering have enabled the observation of nuclear motion in molecules [2-6] and has the potential to track electronic motion [7]. In this scheme, an x-ray probe pulse scatters off an ensemble of optically photoexcited molecules, and this process is repeated for a series of delays between the optical excitation and the x-ray probe. One advantage of x-ray scattering is that the x rays scatter from all of the electrons in the molecular system under study and thus embed spatial information about the electronic charge distribution of a molecule. This has enabled the probing of parallel and perpendicular transitions in N-methylmorpholine [8], the direct observation of bound and dissociative motion in molecular iodine [3,4], the resolving of various molecular trajectories in 1,3-cyclohexadiene [2], and vibrational coherence following electronic relaxation [9].

Time-resolved x-ray scattering, however, is not a direct probe of nuclear position. Since scattering takes place in momentum, or rather reciprocal space, it is necessary to either fit the data to a model [2,5] or invert the data from reciprocal space to real space to recover the nuclear dynamics [3,4]. The inverse problem to obtain the nuclear dynamics, i.e., the pair-distribution function, is difficult because the observed momentum transfer $\vec{Q} = \vec{k}_0 - \vec{k}_s$ between the incoming and outgoing x-ray photon is limited to a few Å⁻¹ in an x-ray freeelectron laser (FEL) [10]. For example, the scattered x rays in a typical experiment at the LINAC Coherent Light Source are restricted between $Q_{\min} = 0.1 - 1$ Å⁻¹ and $Q_{\max} = 2\pi/\lambda \approx$ 4.5 Å⁻¹ [11]. This restricts the spatial resolution to $\Delta R =$ $2\pi/(Q_{\max} - Q_{\min}) > 1.4$ Å. There is however no comparable technical restriction on the pump-probe delay τ , which may range over a long time τ_R with fine steps $\Delta \tau$. A temporal Fourier transform of the scattering pattern $I(x, y, \tau)$ forms the frequency-resolved x-ray scattering (FRXS) signal $\tilde{I}(x, y, \omega)$, where the maximum observable beat frequency is given by $2\pi/\Delta \tau$ and the frequency resolution is $2\pi/\tau_R$. The temporal Fourier transform thus generates a high-resolution representation of TRXS.

Temporal Fourier transform methods have been developed for Fourier transform inelastic x-ray scattering to characterize phonon modes in crystals to arbitrary frequency resolution [12]. Temporal transform methods for TRXS from gases have been used to isolate diatomic vibrations [13].

Here the expression for FRXS of a dissociating diatomic molecule is derived, and the frequency spectrum is shown to isolate dissociative motion along lines in reciprocal space and reciprocal time, Q and ω , despite dissociation not being periodic in real space, i.e., dissociations follow a trajectory like $R(\tau) = R_0 + v\tau$, see Ref. [14]. The analytical results are then confirmed through comparison to measured experimental data, where two dissociative states are observed: one parallel to the pump laser's polarization and one perpendicular.

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FIG. 1. Difference image on the x-ray detector (CSPAD) for a delay of 120 fs after pumping with 520-nm light.

II. EXPERIMENT

The experimental apparatus to study the time-resolved xray scattering of molecular iodine following photoexcitation by 520-nm light has been described elsewhere [3,11,13] and a short description and schematic is provided in Ref. [13]. An attempted real-space reconstruction of the data was published previously [3]. In contrast, the analysis in this paper using FRXS highlights an approach to isolate and characterize dissociations measured using TRXS.

The specifics of the experiment are as follows. A 50-fs, $20-\mu$ J, vertically polarized pulse of 520-nm light is focused into a gas cell containing around 50 Torr of molecular iodine at 100 °C. Following the optical pump, a 50-fs, 2-mJ, horizontally polarized x-ray probe arrives at the scattering cell with variable delay. The resulting scattering is measured in the forward direction by the Cornell-SLAC Pixel Array Detector (CSPAD) [10]. Scattering is measured at each pump-probe delay, resulting in the difference image shown in Fig. 1.

At 520 nm, one photon will excite molecular iodine to the bound $B^{3}\Pi_{0u}^{+}$ state or the dissociative ${}^{1}\Pi_{u}$ state. Two photons can access higher dissociative states, including the ${}^{1}\Sigma_{g}^{+}$ state. The excitation begins near the equilibrium position of the ground $(X^{1}\Sigma_{g}^{+})$ state at $R_{0} = 2.666$ Å. The corresponding potential energy curves are shown in Fig. 2 from Ref. [15]. In the bound B state, 520 ± 5 -nm light excites highly anharmonic vibrations with periods between T = 520and 650 fs and angular frequencies between $\omega = 9.6$ and 11.9 THz. In the dissociative ${}^{1}\Pi_{u}$ state, the internuclear separation increases at a rate of v = 16 Å/ps, and in the dissociative ${}^{1}\Sigma_{g}^{+}$ state, v = 20 Å/ps. The beat frequencies and periods were derived from [16] assuming a pump wavelength of 520 nm, and velocities can be derived from the potential energy curves from [15]. The initial position R_0 , the above frequency ω , and velocities v are observed using FRXS, which will now be described.

III. THEORY

As discussed in previous papers [13,17,18], time-resolved x-ray scattering may be expressed as a product of three factors

$$\frac{dI}{d\Omega} = \frac{d\sigma_{\rm Th}}{d\Omega} I_0 \langle F(\vec{Q}, \tau) \rangle, \tag{1}$$



FIG. 2. Following photoexcitation by 520-nm light, a single photon may excite high in the bound *B* state ($\omega = 9.6-11.9$ THz) or the dissociative ${}^{1}\Pi_{u}$ state (v = 16 Å/ps). Two photons excite the dissociative ${}^{1}\Sigma_{g}^{+}$ state which shares the same symmetry as the ground *X* state (v = 20 Å/ps). These states are identified in the frequency-resolved scattering in Figs. 3(e) and 4(b). The potential energy curves are from Ref. [15].

where $\frac{dI}{d\Omega}$ is the number of photons scattered into a solid angle Ω , $\frac{d\sigma_{\text{Th}}}{d\Omega}$ is the Thomson scattering cross section, I_0 is the incident x-ray intensity, and $\langle F(\vec{Q}, \tau) \rangle$ is a time- and angle-dependent polarization-corrected scattering probability given by

$$\langle F(\vec{Q},\tau)\rangle = 2|f_A(Q)|^2[1+S(\vec{Q},\tau)]$$
 (2)

for a homonuclear diatomic molecule. In Eq. (2), $f_A(Q)$ is the atomic form factor and $S(\vec{Q}, \tau)$ is the molecular scattering factor, which encodes the internuclear separations. This expression is correct in the limit where the independentatom approximation holds, i.e., heavy atoms and insufficient time resolution to observe coherent effects between electronic states [18–21]. Within this approximation $S(\vec{Q}, \tau)$ may be considered for each electronic state independently and may be expressed as

$$S(\vec{Q},\tau) = \int d\vec{R} \,\rho(\vec{R},\tau) \cos(\vec{Q}\cdot\vec{R}),\tag{3}$$

where $\rho(\vec{R}, \tau)$ is the internuclear probability density on some electronic state [17,22].

Turning to FRXS, consider a perfectly aligned classical diatomic molecule, which dissociates along a trajectory $\vec{R}(\tau) = (R_0 + v\tau)\hat{e}_z$ for $\tau > 0$. For this special case, the molecular scattering factor from Eq. (3) evaluates to

$$S(Q_z, \tau) = \cos Q_z R_z(\tau). \tag{4}$$

Now FRXS is defined as

$$\tilde{S}(\vec{Q},\omega) = \int_{-\infty}^{+\infty} d\tau \, e^{-i\omega\tau} S(\vec{Q},\tau).$$
(5)

Therefore, for the above special case in Eq. (4), the FRXS signal is

$$\tilde{S}(Q_z, \omega) = \frac{1}{2} [e^{i\omega R_0/v} \delta(\omega - vQ_z) + e^{-i\omega R_0/v} \delta(\omega + vQ_z)].$$
(6)



FIG. 3. (a) Theoretical time-resolved x-ray scattering following photoabsorption of two 520-nm photons onto the dissociative ${}^{1}\Sigma_{g}{}^{+}$ state, using v = 20 Å/ps and $R_{0} = 2.666$ Å. (b) Power spectrum under similar conditions generated using Eq. (9). (c) Real part of the frequency spectrum of (b). White lines run parallel to $\omega = vQ$ and the circles identify the nodes at $\frac{\omega R_{0}}{v} = \frac{n\pi}{2}$, where n = 3 and 5. Note that a phase shift was applied to Eq. (8) such that (c) reproduces (f). (d) Experimental time-resolved x-ray scattering following photoabsorption of 520-nm light. This is the component of the data projected onto the second-order Legendre polynomial $P_{2}(\cos \theta)$. (e) Power spectrum of (d). The $v = 19.9 \pm 0.2$ Å/ps dissociation is observed and the bound *B* state motion peaks at $\omega = 11.6 \pm 1.1$ THz. (f) Real component of the frequency spectrum.

This equation has two important properties: (i) The maxima of the FRXS lie along $\omega = vQ$ and (ii) the phase evolves like $\phi = \omega R_0/v$. The rate of increase of internuclear separation in a dissociation may then be obtained by fitting the positions of the maxima to a line, and the initial position of the dissociation may be obtained by fitting the phase along those maxima to another line, as will be demonstrated in Sec. IV.

Before turning to the experimental analysis, the physical alignment of a diatomic will now be considered. Following excitation by a polarized laser pulse, the angular distributions will go as $\cos^{2n} \theta$ for parallel transitions or $\sin^{2n} \theta$ for perpendicular transitions, where *n* indicates the number of photons absorbed and θ is the angle with respect to the laser polarization axis. These distributions may be expressed as a linear combination of Legendre polynomials $P_l(\cos \theta)$ such that the molecular scattering factor from Eq. (3) may be rewritten as

$$S(\vec{Q},\tau) = \sum_{l} P_{l}(\cos\theta) S_{l}(Q,\tau), \tag{7}$$

where

$$S_l(Q,\tau) = \int dR R^2 \rho_l(R,\tau) j_l(QR), \qquad (8)$$

with $\rho_l(R, \tau)$ the projection of the nuclear probability function onto a given Legendre polynomial and $j_l(QR)$ the spherical Bessel functions. Now the FRXS may be considered for a given Legendre order $\tilde{S}_l(Q, \omega) = \int_{-\infty}^{+\infty} d\tau \ e^{-i\omega\tau} S_l(Q, \tau)$. As shown in Appendix A, the FRXS for a dissociating diatomic is approximately given by

$$\tilde{S}_{l}(Q,\omega) \approx \frac{e^{i\omega R_{0}/v}}{2iQv} [E_{1}(-i(QR_{0}-\omega R_{0}/v)) - E_{1}(i(QR_{0}+\omega R_{0}/v))]$$
(9)

for each Legendre order, where $E_1(z) = \int_z^{\infty} dt \ e^{-t}/t$ is an exponential integral. To show that Eq. (9) shares the same important features as the first derivation in Eq. (6), the result is shown in Figs. 3(b) and 3(c), which demonstrate that (i) the maxima of the FRXS lie along $\omega = vQ$ and (ii) the phase evolves like $\phi = \omega R_0/v$.



FIG. 4. (a) Experimental time-resolved x-ray scattering following photoabsorption of 520-nm light. This is the isotropic component of the data found by projecting the CSPAD image at each time delay onto the zeroth-order Legendre polynomial $P_0(\cos \theta)$. (b) Power spectrum of (a). There are two dissociations evident in the data: $v = 19.9 \pm 0.2$ Å/ps and $v = 16.4 \pm 0.2$ Å/ps. There is also the bound *B* state motion peaked at $\omega = 11.6 \pm 1.1$ THz.

IV. DISCUSSION

To validate the theory derived above, a comparison to the experimental data is now made. For each pump-probe delay of the TRXS as shown in Fig. 1, the Thomson cross section and atomic form factors are divided out, $I(x, y, \tau) \rightarrow S(x, y, \tau)$, and the pixel coordinates (x, y) are mapped onto the scattering coordinates (Q, θ) using the method described in Appendix B, $S(x, y, \tau) \rightarrow S(Q, \theta, \tau)$. Following the coordinate mapping, $S(Q, \theta, \tau)$ is projected onto the zeroth through tenth Legendre polynomials to obtain $S_l(Q, \tau)$. Figures 3(d) and 4(a) show $S_0(Q, \tau)$ and $S_2(Q, \tau)$, respectively.

These Legendre projections are then used to generate the FRXS, $\tilde{S}_0(Q, \omega)$ and $\tilde{S}_2(Q, \omega)$, through a discrete Fourier transform (DFT). The power spectrum following the DFT is shown in Figs. 3(e) and 4(b). The power spectrum allows for the identification of the bound state and two dissociations. The bound state is peaked at $\omega = 11.6 \pm 1.1$ THz, the first dissociation has a final velocity of 16.4 ± 0.2 Å/ps, and the second dissociation has a final velocity of 19.9 ± 0.2 Å/ps. These results align with the inferred values for the ${}^{1}\Pi_{u}$ and $B{}^{1}\Sigma_{g}{}^{+}$ states, respectively, as derived from Ref. [15]. For information on why the bound state appears at its beat frequency in the power spectrum, see [13] for details on characterizing bound-state motion using FRXS.

The dissociations can also be observed in the original images on the detector following the temporal Fourier transform $\tilde{I}(x, y, \omega)$. The dissociations present themselves as outward moving crescents on the detector image in reciprocal space and reciprocal time. The slower dissociation appears first on the detector perpendicular to the pump polarization as ω is increased, whereas the perpendicular alignment has been seen elsewhere [23,24]. The faster dissociation then appears moving parallel to the pump polarization. This effect is shown in Figs. 5(a) and 5(b) as well as the GIF included in the Supplemental Material [25].

The faster dissociation as seen in the second-order Legendre projection in Fig. 3(e) will be used to demonstrate that FRXS can characterize the dissociation velocity and initial position. To obtain the dissociation velocity, the position of the maximum at each momentum transfer Q is extracted from the power spectrum in Fig. 3(e). The positions of the maxima (Q, ω) are then used to fit the line $\omega = vQ$ as shown in Fig. 6(a). This method obtains a velocity of $v = 19.9 \pm$ 0.2 Å/ps as compared to the predicted 20 Å/ps from the dissociative ${}^{1}\Sigma_{g}^{+}$ state. Now using the measured velocity, the phase along the line $\omega = vQ$ is extracted to find the initial position R_0 . The phase as a function of the angular frequency is fit to $\phi = \omega R_0 / v + \phi_0$ to obtain the initial position as shown in Fig. 6(b). [For reference, the real part of the frequency spectrum is shown in Fig. 3(f).] This method obtains an initial position of $R_0 = 2.3 \pm 0.4$ Å as compared to the known value of 2.666 Å.

V. CONCLUSION

An analysis method for leveraging FRXS to characterize dissociative motion has been derived and applied to experimental data. The obtained values for initial position and dissociation velocities align with the expectations for the ${}^{1}\Pi_{\mu}$ and ${}^{1}\Sigma_{g}^{+}$ states of molecular iodine. The different aspect of this approach is that an interpretable and compact representation of the experimental measurement may be obtained in reciprocal space and reciprocal time without the difficulty of inverting the measurement to the traditional space and time representation. Thus, FRXS presents an alternative to traditional analyses of TRXS. The traditional approach is limited by the range of momentum transfer Q that is accessible at FELs. Frequency-resolved x-ray scattering does not suffer this limitation, and in fact FRXS leverages the strengths of FELs, namely, fine time resolution and fast data accumulation. This enables a long range of pump-probe



FIG. 5. Power spectrum $\tilde{I}(x, y, \omega)$ of the time binned detector images $I(x, y, \tau)$. (a) Power spectrum at $\omega = 20.9$ THz across all pixels. Parallel to the laser field (y axis), we observe the two-photon 20-Å/ps dissociation beginning to emerge from the beam block. Perpendicular to the laser field (x axis) we observe the one-photon 16-Å/ps dissociation. (b) At $\omega = 27.9$ THz, the two-photon dissociation has moved outward on the detector as the frequency was increased, and the one-photon dissociation is no longer visible. In the Supplemental Material [25], a GIF is included showing the two dissociations propagating outward as frequency is increased.

delays to be measured in an experiment, thereby improving the frequency resolution of an experiment while maintaining sufficient temporal resolution to measure high beat frequencies. These advantages have been leveraged to obtain compact representations of dissociations along lines in reciprocal space and reciprocal time, demonstrating an alternative to traditional analyses of time-resolved x-ray scattering for gas-phase photochemistry.

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FIG. 6. (a) Position of the maxima at each momentum transfer Q along the dissociation line in Fig. 3(e) shown in red. The fit to $\omega = vQ$ is shown in black, which finds $v = 19.9 \pm 0.2$ Å/ps, enabling the identification of dissociation along the ${}^{1}\Sigma_{g}{}^{+}$ state. Error bars are the bin size in Q from Fig. 3(e) and error bars in ω are determined by the scan range. (b) Phase along the dissociation line in Fig. 3(f) shown in red. The fit to $\phi = \omega R_0 / v + \phi_0$ is shown in black, which recovers the initial position before dissociation as $R_0 = 2.3 \pm 0.4$ Å. Error bars are the standard deviation of the phase.

APPENDIX A: FRXS OF AN ALIGNED DISTRIBUTION

To reproduce the exact structure of the FRXS given by a dissociation, the alignment of the molecule needs to be considered. For example, an isotropic distribution will project onto the zeroth-order Legendre and have a molecular scattering distribution given by $S_0(Q, \tau) = \int dR R^2 \rho(R, \tau) j_0(QR)$, where $j_0(QR)$ is the zeroth-order spherical Bessel function. A $\cos^2 \theta$ distribution will project onto both the zeroth- and second-order Legendre polynomials and have a molecular scattering distribution with both the $S_0(Q, \tau)$ component as well as $S_2(Q, \tau) = \int dR R^2 \rho(R, \tau) j_2(QR)$, where $j_2(QR)$ is the second-order spherical Bessel function.

For simplicity, consider an aligned distribution of molecules with $R^2\rho(R, \tau) = \delta(R - R(\tau))$, and then $S_l(Q, \tau) = j_l(QR(\tau))$ for $\tau > 0$. The frequency-resolved scattering is accordingly

$$\tilde{S}_l(Q,\omega) = \int_0^\infty d\tau \, e^{-i\omega\tau} j_l(QR(\tau)). \tag{A1}$$

Each even spherical Bessel function contains a $\sin x/x$ term, which by observation makes the largest contribution to this integral. Focusing the derivation to this term, the approximate



FIG. 7. For a linearly polarized pump pulse, the momentum transfer \vec{Q} is decomposed into its projection onto the \hat{e}_y axis and the $\hat{e}_x - \hat{e}_z$ plane. The large circle depicts the Ewald sphere, which represents all possible \vec{Q} for an elastic scattering experiment.

solution can be found by taking

$$\tilde{S}_{2l} \approx \int_0^\infty d\tau \, e^{-i\omega\tau} \frac{\sin QR(\tau)}{QR(\tau)}.$$
 (A2)

Dissociation is the focus here, so $R(\tau) = R_0 + v\tau$. Then taking $u = QR(\tau)$ and expanding sine as a difference of exponentials, the integral becomes

$$\tilde{S}_{2l}(Q,\omega) = \frac{e^{i\omega R_0/v}}{2iQv} \left[\int_{QR_0}^{\infty} \frac{e^{iu(1-\omega/Qv)}}{u} - \int_{QR_0}^{\infty} \frac{e^{-iu(1+\omega/Qv)}}{u} \right].$$
(A3)

This may be identified as the difference of exponential integrals after taking two additional *u* substitutions with $a = u(1 - \omega/Qv)$ and $b = u(1 + \omega/Qv)$ such that

$$\tilde{S}_{2l}(Q,\omega) = \frac{e^{i\omega R_0/v}}{2iQv} \left[\int_{QR_0-\omega R_0/v}^{\infty} da \frac{e^{ia}}{a} - \int_{QR_0+\omega R_0/v}^{\infty} db \frac{e^{-ib}}{b} \right]$$
$$= \frac{e^{i\omega R_0/v}}{2iQv} [E_1(-i(QR_0-\omega R_0/v)) - E_1(i(QR_0+\omega R_0/v))], \quad (A4)$$

where $E_1(z) = \int_z^{\infty} dt \ e^{-t}/t$ is an exponential integral. Equation (4) shares the same features described in the simplified derivation in the text for perfectly aligned diatomics: (i) The maxima of the FRXS lie along $\omega = Qv$ and (ii) the phase evolves like $\phi = \omega R_0/v$.

APPENDIX B: COORDINATE MAPPING

For a diatomic excited along a polarized laser pulse, the relevant angular decomposition (θ, ϕ) is shown in Ref. [13]. Mapping from the detector image I(x, y) set a distance L away from the scattering center onto the molecular frame $I(Q, \theta)$, as sketched in Fig. 7, is a simple geometry problem. First, the momentum transfer must be decomposed in the directions

parallel to the field, \hat{e}_y , and perpendicular to the field, the $\hat{e}_x \cdot \hat{e}_z$ plane. The parallel decomposition is given by $Q_y = \frac{k_{0y}}{\sqrt{R^2 + L^2}}$, where $R = \sqrt{x^2 + y^2}$ is the distance from the center of the detector, and the perpendicular decomposition is given by $Q_{\perp} = k_0 \sqrt{\frac{x^2 + L^2}{R^2 + L^2} + 1 - \frac{2L}{\sqrt{R^2 + L^2}}}$. With that decomposition in hand, (Q, θ) is determined by $Q^2 = Q_y^2 + Q_{\perp}^2$ and $\tan \theta = Q_{\perp}/Q_y$.

APPENDIX C: ERROR PROPAGATION AND DATA ANALYSIS

For each shot the pump-probe delay τ'_i is measured and then binned into some time bin $\tau_j \pm \Delta \tau$. This allows for the generation of the mean scattered intensity at each time delay

$$I(x, y, \tau_j) = \frac{1}{N_j} \sum_{i} I(x, y, \tau'_i),$$
 (C1)

where (x, y) indicate each pixel on the CSPAD detector. From these images, the unpumped signal is subtracted to find the difference scattering $\Delta I(x, y, \tau_j) = I(x, y, \tau_j) - I_u(x, y)$.

The difference images are then divided by the Thompson cross section $\frac{d\sigma}{d\Omega}$ [26], iodine's atomic form factor $|f_I(Q)|^2$ [22], and the correction factor for attenuation in the scattering cell [11]. This results in the difference molecular scattering factor $\Delta S(x, y, \tau_j)$ up to an overall factor of the x-ray intensity. Now the variation at each pixel position is generated by

$$\sigma^{2}(x, y, \tau_{j}) = \operatorname{Var}[\Delta S(x, y, \tau_{j})]$$
$$= \frac{1}{N_{j}} \sum_{i} |\Delta S(x, y, \tau_{i}') - \Delta S(x, y, \tau_{j})|^{2}. \quad (C2)$$

The variation is then propagated through the analysis as follows.

For the projection of $\Delta S(x, y, \tau'_i)$ onto Legendre polynomials, the coordinates are first mapped from (x, y) onto (Q, θ) as described in Appendix B. Then mapping $\Delta S(Q, \theta, \tau)$ onto the Legendre coefficients $\Delta S_l(Q, \tau)$ is achieved through a χ^2 minimization. The χ^2 model is defined by

$$\chi^2 = \sum_i \frac{(p_i - s_i)^2}{\sigma_i^2},$$
 (C3)

where p_i is the fitted function, s_i is the data, and σ_i^2 is the variance. For a linear model, the fitted function may be expressed as $p_i = \sum_j x_j f_j(Q_i)$, where x_j are the model coefficients. Then the solution to the χ^2 minimization is

$$x_k = \sum_i A_{ki} s_i / \sigma_i^2, \tag{C4}$$

where $A = (f^T \sigma^{-2} f)^{-1} f$. The associated error for the solution x_k is then

$$\sigma_k = \sqrt{\sum_j \left(A_{kj}\sigma_j^{-1}\right)^2},\tag{C5}$$

as shown in [27]. For the Legendre projection of $\Delta S(Q, \theta, \tau)$ onto $\Delta S_l(Q, \tau)$, the fitted function is $f_j(\theta_i) = P_j(\cos \theta_i)$, where j = 0, 2, ..., 10 are used. For the temporal Fourier transform of the Legendre coefficients $\Delta S_l(Q, \tau)$, a discrete Fourier transform is used. As the Fourier transform is a unitary transform, the standard deviation for each frequency element is simply the sum in

quadrature of the errors $\sigma_l(Q, \tau_i)$, where

$$\sigma_l(Q, \omega_j) = \sqrt{\sum_j \sigma_l^2(Q, \tau_j)}.$$
 (C6)

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