Quantum coherence in photo-ionisation with tailored XUV pulses

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Abstract
Ionisation with ultrashort pulses in the extreme ultraviolet (XUV) regime can be used to prepare an ion in a superposition of spin–orbit substates. In this work, we study the coherence properties of such a superposition, created by ionising xenon atoms using two phase-locked XUV pulses at different frequencies. In general, if the duration of the driving pulse exceeds the quantum beat period, dephasing will occur. If however, the frequency difference of the two pulses matches the spin–orbit splitting, the coherence can be efficiently increased and dephasing does not occur.

Keywords: coherent ionisation, atomic double slit, ultra-fast entanglement

(Some figures may appear in colour only in the online journal)

1. Introduction
The wave nature of matter is central to the quantum mechanical description of the microcosmos; therefore coherence—a measure of the ability to produce stationary interference patterns—is an important property of any quantum system. An example of a coherent system is the superposition of two pure states, \(|\psi\rangle = a|1\rangle + b|2\rangle\); such superpositions form the basis for the field of quantum information, where they are used to represent qubits. The manipulation of qubits for quantum computing necessarily requires that the coherence of the system is retained; if not, the information contained within the qubit is lost. In quantum optics, superpositions between two states may be created via a transition between the two states with an appropriately tailored pulse (e.g. a \(\pi/2\)-pulse) (figure 1(a)).

Superpositions of states can also be achieved by direct excitation using short light pulses (figure 1(b)), provided the bandwidth of the pulse is larger than the energy difference \((\omega_{21})\) between the two states. This requires a pulse duration short enough, \(\tau \lesssim 2\pi/\omega_{21}\). If the superposition is successfully created, it may be observed through quantum beats (Salour and Cohen-Tannoudji 1977, Teets et al 1977, Mauritsson et al 2010, Tzallas et al 2011) which usually last substantially longer than the pulse duration. The characteristic decay time is termed the coherence time. In the cases depicted in 1(a) and (b), the light couples the bound states and enables coherent population transfer.

Another way to produce a superposition of states is via short-pulse ionisation, when the ion is left in a coherent superposition of final states, e.g. due to spin–orbit interaction. This can be done using either high-frequency (figure 1(c)) or high-intensity short-pulse (figure 1(d)) radiation. As previously, the bandwidth of the ionising pulse has to exceed the energy splitting between the ion states. Kurka et al (2009) investigated case 1(c) by photo-ionising neon using short XUV pulses from a free-electron laser. A coherent superposition of the ionic fine-structure substates was prepared by photo-ionising neon using short XUV pulses from a free-electron laser. A coherent superposition of the ionic fine-structure substates was prepared and probed by subsequent ionisation. Using a strong laser field (figure 1(d)), Goulielmakis et al (2010) photo-ionised krypton, leaving the residual Kr+ ion in a coherent superposition of the ionic substates. The quantum beat was observed by probing with a delayed attosecond (as) XUV pulse, as long as the duration of the ionising pulse was shorter than the
quantum information; generation harmonics such as those resulting from high-order harmonic fields, maximising the coherence between states, independent of the driving field, providing a new ionisation scheme to control and steer the ionisation process in such a way that the quantum paths always overlap speculatively, allowing the coherence of superpositions of states produced either directly by photo-excitation (figure 1(b); Tzallas et al. 2011, Klundert et al. 2013), single-photon ionisation (figure 1(c); Nikolopoulos 2013), or strong-field ionisation (figure 1(d); Pabst et al. 2011, 2016).

As mentioned above, the creation of a coherent superposition in the cases depicted in figures 1(b)–(d), requires sufficiently high bandwidth of the exciting/ionising radiation. As the pulse duration increases, or the energy separation between electronic states increases, states become spectrally resolvable. Excitation/ionisation to one state or the other can then be seen as distinguishable quantum paths taken by the system (see figure 2). When the spectral overlap between these quantum paths decreases, the coherence between states diminishes. While a spectral representation provides meaningful physical insight, it does not allow understanding how coherence is built up in real time. Therefore, a temporal representation, based on an atomic clock constituted by the quantum beat period, is very useful to determine whether decoherence will occur or not during light–matter interaction. As long as the pulse duration is shorter than a quantum beat period, the atomic clock will not dephase, since there is still appreciable overlap between the quantum paths (see figure 2). For ionisation of noble gases, the available quantum beat periods span an order of magnitude (see table 1). However, ultrashort pulses are still necessary to manipulate the coherence.

In this article, we present a theoretical study of single-photon ionisation of xenon atoms using XUV pulses, tailored in such a way that the quantum paths always overlap spectrally, providing a new ionisation scheme to control and maximise the coherence between states, independent of the pulse duration. This is achieved by employing a bichromatic or multi-colour ionising field, consisting of phase-locked harmonics such as those resulting from high-order harmonic generation (HHG), providing the spectral components fulfilling a certain resonance condition. We investigate the tolerance of this resonance condition, i.e. how strict the requirements on the driving field are, with respect to the excitation frequencies, pulse duration, and temporal structure, for maintaining a certain level of coherence. We find the existence of resonant conditions, which correspond to a situation where multiple quantum paths lead to the same photo-electron state.

The paper is organised as follows; in the following section, we present our numerical calculations based on the fully correlated time-dependent Schrödinger equation (TDSE) in the case of weak-field ionisation, as well as the theoretical tools that are used to calculate the evolution of the superposition of states in the presence of the driving field. We also introduce the density matrix formalism used to analyse the coherence of the quantum system. Atomic units are used throughout, unless otherwise stated. Based on the numerical calculations, in section 3, we study ionisation using two harmonic components, and investigate the relation between their energy separation and the spin–orbit splitting using a temporal model of the ionisation dynamics. From this, we extract a generalised quantum beat condition. Finally in section 4, we derive a spectral model for the general case of ionisation by multi-colour fields. This enables us to capture the essential physics of the observed phenomena. We conclude with a short discussion about the foundation of the work in relation to quantum mechanics, suggesting that coherence between states should exist as long as ionisation pathways are indistinguishable by the measurement.

### Table 1. Some properties of heavy noble gases. \( \Delta E_{\text{so}} \) is the spin–orbit splitting of the ionic ground state \( n\pi^2 \Sigma^+ \). The quantum beat period, \( T = 2\pi \Delta E_{\text{so}}^{-1} \), is the intrinsic atomic clock associated with two states separated by an energy difference of \( \Delta E_{\text{so}} \).

| Element | Z | n | \( \Delta E_{\text{so}} \) (eV) | \( T \) (fs) |
|---------|---|---|-----------------|--------|
| Ne      | 10| 2 | 0.096760 24     | 42.8   |
| Ar      | 18| 3 | 0.177493 68     | 23.3   |
| Kr      | 36| 4 | 0.665 808       | 6.2    |
| Xe      | 54| 5 | 1.306 423       | 3.2    |

### Figures

**Figure 1.** Different ways of preparing coherent superpositions using light; (a) excitation of a two-level system, such as those used for qubits in quantum information; (b) coherent excitation from the ground state to two excited bound states; (c) single-photon ionisation, with the ion left in a superposition of substates; (d) strong-field ionisation, also leaving the ion in a superposition of substates.

**Figure 2.** Overlap (shaded area) between two quantum paths (1 and 2) separated by \( \Delta E \). The spectral bandwidth, \( \Omega \), of the ionising pulse is inversely proportional to the duration of the pulse; increasing the pulse duration thus leads to decreased overlap between the quantum paths.
is the atomic Hamiltonian, $E$ the quantum state of the photo-electron, kinetic energy, which depends on the four pathways to the continuum. If $\omega_0 \approx \Delta E_{s-o}$, two of the quantum paths (absorption of $\Omega_>$ and the ion in $2P_{1/2}$; absorption of $\Omega_<$ and the ion in $2P_{3/2}$) lead to the same photo-electron energy.

**Figure 3.** Schematic energy diagram of a noble gas (heavier than He, i.e. with a spin–orbit splitting of the first ionic ground state $np^5 \, 2P_{3/2}$, $j_i = 3/2$, 1/2) photo-ionised with a tailored XUV pulse consisting of two frequencies with $\Omega_+ - \Omega_- = \omega_0$. The energy scale is that of the photo-electron kinetic energy, which depends on the difference of two frequencies with $\omega_0$, which in xenon is 1.3 eV. We ionise with a weak XUV pulse leading to an ion in either $2P_{3/2}^{0}$ or $2P_{1/2}^{1}$, resulting in four different pathways. If the frequency difference is equal to the spin–orbit spacing, there will be two (indistinguishable) pathways to the same final photo-electron energy; we call this the resonant case. We introduce the detuning ratio $d \equiv \omega_0 / \Delta E_{s-o}$, and study photoionization in the vicinity of this resonance ($d \approx 1$).

The calculations are performed by solving the TDSE in a limited subspace,

$$i\partial_t |\psi(t)\rangle = \mathcal{H}(t)|\psi(t)\rangle,$$

where the Hamiltonian in the dipole approximation is

$$\mathcal{H}(t) = \mathcal{H}_0 + \mathcal{E}(t)z,$$

$\mathcal{H}_0$ is the atomic Hamiltonian, $\mathcal{E}(t)$ the electric field, and $z$ is the dipole operator for linear polarisation along the z axis. The solution is found by propagating the initial state (the neutral ground state) to time $t$

$$|\psi(t)\rangle = \mathcal{U}(t, 0)|\psi_0\rangle,$$

where the short-time propagator $\mathcal{U}(t + \Delta t, t)$ is approximated by a Magnus (1954) propagator of fourth order (Saad 1992, Alvermann et al 2012).

The time-dependent wavefunction is expanded as

$$|\psi(t)\rangle = c_0(t)|\psi_0\rangle + \sum \sum \int d\epsilon \, c_{i \ell}^*(t; \epsilon)|\epsilon \ell\rangle,$$

where $|\psi_0\rangle$ is the ground state $nsnp^6\, 1S_0$ with energy $-I_p$, $c_0(t)$ the complex, time-dependent amplitude, $i$ denotes the final state of the ion, and $|\epsilon \ell\rangle$ the quantum state of the photo-electron with angular momentum $\ell$ and energy $\epsilon$ (related to the momentum $k$ by $\epsilon = k^2/2$). The ionisation channels formed by different possible combinations of $i$ and $\ell$, are listed in table 2, in the case of $jK$ coupling. $jK$ (or pair) coupling is defined as (Cowan 1981) $j_i + \ell = K$ and $K + s = J$, where $j_i$ is the total angular momentum of the parent ion, which couples to the angular momentum of the electron $\ell$ to form an intermediate $K$. The levels are then written as $\gamma_5^{(2S+1)J_i}k\ell 2S+1|K_j\rangle$, where $\gamma_5$ is the electron configuration of the ion.

The ansatz (4) turns the TDSE (1) into a set of coupled ordinary differential equations (ODE):

$$i\partial_t \epsilon(t) = \mathcal{H}(t) \epsilon(t),$$

where the vector $\epsilon(t)$ consists of the expansion coefficients in (4), and the Hamiltonian matrix is given by

$$\mathcal{H}(t) = -|\psi_0\rangle I_p|\psi_0\rangle + \sum \int d\epsilon \, |\epsilon \ell\rangle \epsilon (|\epsilon \ell\rangle$$

$$+ \mathcal{E}(t) \sum \int d\epsilon \, |\epsilon \ell\rangle \langle \epsilon \ell|z|\psi_0\rangle$$

$$+ \sum \int d\epsilon \, |\epsilon \ell\rangle \langle \epsilon \ell|z|\epsilon' \ell'\rangle \langle \epsilon' \ell'| + c.c. \right].$$

In the field-free basis, $\mathcal{H}_0$ is simply a diagonal matrix, with the energies of the photo-electron with respect to the lowest ionisation threshold as matrix elements. The interaction term couples the ground state to the continuum and the continua to each other. In the weak-field limit, however, the partial-wave

| # | Channel configuration |
|---|---|
| 1 | $np^5(2P_{3/2})d k\ell 2(1/2)_{\ell}h$ |
| 2 | $np^5(2P_{3/2})k s\ell 2(3/2)_{\ell}h$ |
| 3 | $np^5(2P_{3/2})k d\ell 2(3/2)_{\ell}h$ |
| 4 | $np^5(2P_{1/2})k s\ell 2(1/2)_{\ell}h$ |
| 5 | $np^5(2P_{1/2})k d\ell 2(3/2)_{\ell}h$ |
| 6 | $nsnp^6(1S_0)k p\ell 2(1/2)_{\ell}h$ |
| 7 | $nsnp^6(1S_0)k d\ell 2(3/2)_{\ell}h$ |

which have a spin–orbit splitting of the ground state ($np^5 \, 2P_{3/2}^0$, $j_i = 3/2$, 1/2), in particular xenon ($n = 5$).

Figure 3 shows a simplified diagram of photo-ionisation of a $np$ electron. The ionic ground state has a spin–orbit splitting, which in xenon is 1.3 eV. We ionise with a weak XUV pulse with two frequency components, whose difference is $\omega_0$. Absorption of the two frequency components, $\Omega_>$ and $\Omega_<$, leads to another in either $2P_{3/2}^{0}$ or $2P_{1/2}^{1}$, resulting in four different pathways. If the frequency difference is equal to the spin–orbit spacing, there will be two (indistinguishable) pathways to the same final photo-electron energy; we call this the resonant case. We introduce the detuning ratio $d \equiv \omega_0 / \Delta E_{s-o}$, and study photoionization in the vicinity of this resonance ($d \approx 1$).
expansion is restricted to total angular momentum \( J \leq 1 \), i.e. no multi-photon processes are considered. Furthermore, ionisation is only allowed from the outer np shell (photoelectron energies in the range 0–11 eV in the case of xenon), to avoid autoionization of embedded Rydberg states in the vicinity of the nsnp \(^{2}S_{1/2}\) threshold (that is, channels 6 and 7 in table 2 need not be considered). We also neglect mixing of singlet and triplet terms. Thus, the only non-zero matrix elements of the dipole operator are \( \langle |ε\ell z| |Ψ_n\rangle \) (and the complex conjugate). The basis functions \( \langle |Ψ_n\rangle \) and \( \langle |ε\ell \ell\rangle \) and the dipole matrix elements \( \langle |ε\ell z| |Ψ_n\rangle \) are obtained using ATSP2K (multi-configurational Hartree–Fock; Froese Fischer et al. 2007) and BSR (close-coupling \( R \)-matrix approach; Zatsarinny 2006, Zatsarinny and Froese Fischer 2009). The dipole matrix elements are spin-averaged by BSR.

The analysis of the coherence is made using the density matrix formalism (Landau and Lifshitz 1977, section 14), where the full density matrix operator is formed from the wavefunction \( |Ψ(t)\rangle \) obtained by solving (1) (time dependence \( t \) omitted below, for brevity)

\[
ρ_T = |Ψ\rangle \langle Ψ|, \quad (5)
\]

with matrix elements of the continuum block

\[
ρ^{ Eld}_{n\ell\ell}(ε_1, ε_2) \equiv c^*_n(ε_1)c_n(ε_2). \quad (6)
\]

We reduce this density matrix to an ion-channel density matrix by first taking the trace over the photo-electron energy \( ε \):

\[
ρ^{ Eld}_{n\ell\ell} \equiv \int dε \langle ε|Ψ\rangle \langle Ψ|ε\rangle
= \int dε \sum_{n,\ell} \sum_{\ell'} \int dε_1 dε_2 \langle ε|1\ell\ell_1\rangle c^*_n(ε_1)c_n(ε_2) \langle i_2\ell_2\ell_2\rangle
= \sum_{n,\ell} \int dε \langle 1|\ell\ell\rangle ρ^{ Eld}_{n\ell\ell}(ε, ε) \langle i_2\ell_2\rangle. \quad (7)
\]

Finally, we construct the ion density matrix by tracing over the photo-electron angular momenta:

\[
ρ_{n\ell\ell} \equiv \sum_{ℓ} \sum_{\ell} \sum_{\ell'} \langle ℓ|1\ell\ell_1\rangle ρ^{ Eld}_{n\ell\ell}(i_2\ell_2\ell_2)\langle ℓ'\ell\ell_2\rangle
= \sum_{ℓ} \sum_{i_2} \langle ℓ|1\ell\ell_1\rangle ρ^{ Eld}_{n\ell\ell}(i_2). \quad (8)
\]

The diagonal elements \( ρ_{namm} \) of this matrix are the populations in each of the ionic states, while the off-diagonal elements \( ρ_{nm\ell\ell} \) contain the coherences between the ionic states. The only non-zero off-diagonal elements are those corresponding to channels for which all quantum numbers are the same (except for the angular momentum of the ion); i.e. only \( ρ_{33} = ρ_{33} \neq 0 \). Decoherence due to decay (through dipole-forbidden interaction) from the upper ionic state to the lower, is neglected.

### 3. The bichromatic case

We investigate the real-time build-up of coherence with a XUV pulse of short or long duration in the non-resonant case (figure 4, left) and the resonant case (figure 4, right). The electromagnetic fields are presented in the upper panels. In both cases, they consist of harmonics 13 and 14 of a fundamental driving field, and a peak intensity of \( 10^8 \text{ W cm}^{-2} \). The short-pulse duration is 500 as, while the long-pulse duration is 15 fs resulting in the formation of a periodic beating of the XUV pulse. Regardless of the pulse duration, the population in the ionic substates (middle panel, shown in solid and dashed lines) increases as the pulse ionises the atom. The population is proportional to the integral of the pulse intensity, hence the appearance of steps in the population build-up.

#### 3.1. The non-resonant case

We first consider the non-resonant case (left panels of figure 4), where the fundamental driving frequency is \( ω_0 = 1.3ΔE_{n,0} \). For a short-pulse duration, the coherence increases during the interaction and stays constant after the pulse has passed. In contrast, for long-pulse duration, the coherence first builds up, and then vanishes at the end of the pulse. The decoherence time, i.e. the time from the onset of the pulse to the decrease of the coherence (see lower panel of figure 4), is approximately equal to the quantum beat period \( T = 2π/ΔE_{n,0} \) of the ionic substates; for xenon with a spin-orbit splitting of 1.3 eV, it is 3.2 fs. This is similar to what was observed by Goulielmakis et al (2010), where the interference pattern, present in the transient absorption signal, disappeared when the ionising infrared pulse exceeded the quantum beat period of 6.2 fs of krypton.

#### 3.2. The resonant case: a generalised quantum beat condition

In the resonant case (right panels of figure 4), the situation is completely different. The coherence is built up during all the interaction time, and remains after the end of the pulse. We can understand the existence of this resonance condition, using a simple two-excited-state model. The two excited states (labelled \( |1\rangle \) and \( |2\rangle \), for simplicity) correspond to the ionic substates, the populations of which result from a periodic sequence of ionisation events (occurring at times \( t_k \) , \( k \in \mathbb{N} \) from the ground state (labelled \( |0\rangle \)). We express this model using an inhomogeneous TDSE for the excited superposition \( |Ψ(t)\rangle = c_1(t)|1\rangle + c_2(t)|2\rangle \), where the ground state constitutes a source term:

\[
i\partial_t |Ψ(t)\rangle = H_0|Ψ(t)\rangle + \tilde{E}(t)(z_{00}|1\rangle + z_{02}|2\rangle),
H_0 \equiv E_1|1\rangle \langle 1| + E_2|2\rangle \langle 2|, \quad z_{00} \equiv \langle i|z|0\rangle. \quad (9)
\]

Assuming no initial excited population, the solution of (9) is given by

\[
|Ψ(t)\rangle = \exp(iH_0t) \left\{ \int d\tau \tilde{E}(\tau')|e^{-iE_1\tau'}z_{00}|1\rangle
+ e^{-iE_2\tau'}z_{02}|2\rangle \right\}. \quad (10)
\]

If we assume that \( \tilde{E}(t) \) is a train of pulses, separated by \( Δt \equiv t_k - t_{k-1} \), we can write the field as

\[
\tilde{E}(t) = \tilde{E}(t) \left\{ \sum_k \delta(t_k)e^{ip_{\text{XUV}}(t_k)} \right\}.
\]
where $\tilde{E}(t)$ is a slowly varying envelope. The solution to the two-excited-state system in this case is

$$\mathcal{M}(t) = \exp(\mathbf{i} \mathbf{H}_{\text{XUV}} t) \sum_k \tilde{E}(t_k) e^{i \mathbf{H}_{\text{XUV}} t_k} [e^{-i \mathbf{H}_{\text{XUV}} t_0} |1\rangle + e^{-i \mathbf{H}_{\text{XUV}} t_0} |2\rangle]
= \sum_k \tilde{E}(t_k) e^{i \mathbf{H}_{\text{XUV}} t_k} [e^{-i \mathbf{H}_{\text{XUV}} t_0} |1\rangle + e^{-i \mathbf{H}_{\text{XUV}} t_0} |2\rangle]
= \sum_k \tilde{E}(t_k) e^{i \mathbf{H}_{\text{XUV}} (t_k-t_0)} |1\rangle + e^{-i \mathbf{H}_{\text{XUV}} (t_k-t_0)} |2\rangle,
$$

where $\tilde{E} \equiv \langle 2|z|0\rangle/\langle 1|z|0\rangle$ is the relative dipole matrix element independent of the instant of ionisation, and—assuming ionisation solely into an unstructured continuum—independent of final electron energy and $\Delta E \equiv E_2 - E_1$, as well.

For the two states to remain coherent, we require that no dephasing is introduced by pulses in the train. Since subsequent pulses are separated in time by $\Delta t$, this is equivalent to requiring that the phase argument in (11) fulfills

$$\Delta E (t_k - t_0) = \Delta E \Delta t (k - a) = 2\pi, \ k, a \in \mathbb{N},$$

is fulfilled, for any integer $r$. $\Delta t$ is a multiple of the quantum beat period for an energy separation $\Delta E$, which does not depend on the duration of the XUV pulse. In the spectral domain, this corresponds to requiring the final electron kinetic energy to be the same.

### 3.3. Maximum coherence achievable: degree of coherence

As seen in figure 4, given that the resonance condition $d = 1$ is fulfilled, the coherence between the ionisation channels seem to increase as long as the pulse is of appreciable amplitude. What is the maximum coherence achievable using this scheme? The theoretical maximum coherence $|\rho_{\text{max}}| = |c_m e^{i \phi_f}|$, is bounded by the Cauchy–Schwartz inequality:

$$|\rho_{\text{max}}|^2 \leq \rho_{\text{min}} \rho_{\text{min}}^*,$$

where $\rho_{\text{min}}$ is the probability of finding the system in state $|\rangle$, which obviously cannot exceed unity. From (12), it is natural to introduce the degree of coherence:

$$\rho_{\text{max}} = \frac{|\rho_{\text{max}}|}{\sqrt{\rho_{\text{min}} \rho_{\text{min}}^*}},$$

which normalises the coherence between two ions to their respective populations. This quantity is useful since, even though the populations in two states are minuscule (as is the case in figure 4) and thereby also the absolute coherence, they may be fully coherent with respect to each other. If this is the case, the degree of coherence will be unity.

Figure 5 shows the degree of coherence, as a function of XUV pulse duration and detuning ratio. For short pulse durations, this quantity is larger than 1/2 (left panel of figure 5). In this regime, the interaction with the XUV pulse occurs within one quantum beat period (3.2 fs), and the four pathways into the continuum (indicated in figure 3) have a
partial spectral overlap. For larger pulse durations, two of the pathways become distinguishable, and do not contribute to the coherence between the ionic substates. The two remaining pathways, namely via $W_<$ leaving the ion in $^3P_{3/2}^o$, and via $W>$ leaving the ion in $^3P_{1/2}^o$, cannot be distinguished when measuring the photo-electron. Provided the resonance condition $d_1 = 1$ is met, the maximum degree of coherence is $1/2$.

Complete decoherence always occurs in the long-pulse limit if $d_1 = 1$ since the quantum pathways are distinguishable. As we will see below, the maximum degree of coherence can be increased by adding more colours to the ionising field.

4. The multi-colour case—ionisation by an attosecond pulse train

We consider now the effect of ionisation with an attosecond pulse train, by including additional harmonic components. To focus on this aspect of the problem, we use a simplified model, where the dipole matrix elements for ionisation are replaced with Heaviside functions (this is an approximation of a flat continuum, i.e. no resonances present):

$$z(\varepsilon) = \theta(\varepsilon),$$

where $\varepsilon$, as before, is the energy of the continuum electron. We still use the ansatz (4), albeit with a compact notation only considering different channels $n$:

$$\Psi(t) = c_0(t) |\Psi_0\rangle + \sum_n \int d\varepsilon \, c_n(t; \varepsilon) |n\rangle.$$  \hspace{1cm} (15)

Inserting this in the Schrödinger equation and applying first-order time-dependent perturbation theory (i.e. the ground state is unaffected by the weak-field ionisation; $c_0(t) = 1$), the solution reads

$$c_n(t) = -i\theta(\varepsilon) \int_{-\infty}^t dt' \, \mathcal{E}(t') \exp(iE_n t').$$ \hspace{1cm} (16)

Evaluating at the time of measurement ($t = +\infty$), we see that the coefficient becomes the Fourier transform of the driving field, evaluated at $-E_n - \varepsilon$. From this, we get the coherence between two channels ($m, n$) as

$$\rho_{mn} = -\int d\varepsilon \, \theta(\varepsilon) \hat{\mathcal{E}}^*_m(E_m + \varepsilon) \hat{\mathcal{E}}(-E_n - \varepsilon),$$ \hspace{1cm} (17)

where $\hat{\mathcal{E}}(\omega)$ designates the Fourier transform of $\mathcal{E}(t)$. This expression shows that the coherence primarily arises from a correlation of the field with itself shifted by the energy difference $\Delta E = E_m - E_n$.

In the long-pulse limit, the spectral components of $\hat{\mathcal{E}}(\omega)$ will become very narrow. If the ionising field is an attosecond pulse train, consisting of the $n_q = q_2 - q_1 + 1$ successive harmonic orders $q \in \{q_1, \ldots, q_2\}$, the shift has to be precisely an integer amount of photons ($\Delta E = N\omega_0$) for the integrand of

Figure 5. Degree of coherence as a function of pulse duration (FWHM of temporal intensity profile) and detuning ratio $d$. The hyperbolas in the main panel indicate the ‘coherence bandwidth’ within which $\hat{\rho}_{SS} > (2|\mathcal{E}|^2)^{-1}$. The black horizontal/vertical lines mark lineouts at constant duration/detuning ratio, which are shown in the lower/left panels. As the pulse duration exceeds the quantum beat period, the peaks become spectrally resolvable, and the total overlap at the central frequency becomes half of that at vanishing pulse duration. The peaks appearing below $d \approx 0.9$ occur when the lower harmonic excites autoionizing Rydberg states between $^3P_{3/2}$ and $^3P_{1/2}$ thresholds.
(17) to be non-zero. Assuming \( q_1 \omega_q, q_2 \omega_q > E_m, E_n \), i.e. all constituent harmonic orders reach above both ionisation thresholds, we have in total \( 2n_q \) pathways into the continuum. In the case of an integer photon shift, \( nN \) of these pathways will overlap, which means

\[
\tilde{\rho}_{mn} = \frac{n_q - N}{n_q}.
\]  

(18)

From (18), we see that by adding more and more colours, we can increase the degree of coherence towards unity. This is illustrated in figure 6 for the case of five harmonics; the maximum degree of coherence is indeed \( 4/5 \), which occurs at \( d = 1 \).

Figure 6 also serves the purpose of illustrating the generalisation of the quantum beat condition (11); e.g. the case \( r = 2 \) corresponds in the time domain to ionising pulses arriving every other quantum beat period. Such a field can be realised through red–blue HHG (resulting in odd and even harmonic orders) from a fundamental frequency \( \Delta E_{x-o}/2 \) (for Xe with \( \Delta E_{x-o} \approx 1.3 \) eV, a fundamental driving wavelength \( \lambda \approx 1.9 \) \( \mu \)m would thus be necessary). From this we see that in the spectral domain, the generalised quantum beat condition (11) is simply \( d = 1/r, r \in \mathbb{Z} \).

Finally, we have also checked that in the long-pulse limit, the case of \( d = 1/2 \), using only odd-order harmonics is no different from the case of \( d = 1 \), using even- and odd-order harmonics. Thus the change of parity (phase difference) of consecutive pulses in the attosecond pulse train has no impact on the coherence, and we conclude that the degree of coherence is essentially dictated by the amount of indiscernible pathways.

5. Conclusion

In summary, we have shown that it is possible to induce coherence between two ionic substates using pulses of duration longer than the quantum beat time of their superposition. This is possible, provided a resonance condition is fulfilled, namely that the driving field has at least two frequency components spaced by precisely the energy difference of the levels of interest. This result shows that when the electron wave packets arising from different pathways have the same kinetic energy, we cannot know which way the ionisation occurred. This situation is reminiscent of a Young’s double slit experiment (see for instance Arndt et al 2005), with the two harmonic orders playing the roles of the two slits. It has to be noted, however, that no interference can be detected in the photo-electron signal, unless the ions are brought to the same final state via some mechanism (e.g. Goulielmakis et al 2010 did this by further exciting from the fine-structure superposition using an XUV pulse). Otherwise, it would be possible to detect the ions and the photo-electrons in coincidence mode and establishing the ionisation pathway.

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