



**HAL**  
open science

## Attosecond spectral singularities in solid-state high-harmonic generation

Ayelet Julie Uzan, Gal Orenstein, Álvaro Jiménez-Galán, Chris Mcdonald,  
Rui Silva, Barry Bruner, Nikolai Klimkin, Valérie Blanchet, Talya  
Arusi-Parpar, Michael Krüger, et al.

► **To cite this version:**

Ayelet Julie Uzan, Gal Orenstein, Álvaro Jiménez-Galán, Chris Mcdonald, Rui Silva, et al.. Attosecond spectral singularities in solid-state high-harmonic generation. *Nature Photonics*, 2020, 14 (3), pp.183-187. 10.1038/s41566-019-0574-4 . hal-03005108

**HAL Id: hal-03005108**

**<https://cnrs.hal.science/hal-03005108>**

Submitted on 18 Nov 2020

**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.

1 **Attosecond spectral singularities in solid-state**  
2 **high-harmonic generation**

3 Ayelet Julie Uzan<sup>\*1</sup>, Gal Orenstein<sup>\*1</sup>, Álvaro Jiménez-Galán<sup>2</sup>,  
4 Chris McDonald<sup>3</sup>, Rui E.F Silva<sup>4</sup>, Barry D. Bruner<sup>1</sup>, Nikolai D.  
5 Klimkin<sup>5,6</sup>, Valerie Blanchet<sup>7</sup>, Talya Arusi-Parpar<sup>1</sup>, Michael  
6 Krüger<sup>1</sup>, Alexey N. Rubtsov<sup>5,6</sup>, Olga Smirnova<sup>2,8</sup>, Misha  
7 Ivanov<sup>2,9,10</sup>, Binghai Yan<sup>1</sup>, Thomas Brabec<sup>3</sup> and Nirit Dudovich<sup>1</sup>

8 <sup>1</sup>Department of Complex Systems, Weizmann Institute of Science, 76100, Rehovot, Israel

9 <sup>2</sup>Max-Born-Institut, Max-Born Strasse 2A, D-12489 Berlin, Germany

10 <sup>3</sup>Department of Physics, University of Ottawa, Ottawa, Ontario K1N 6N5, Canada

11 <sup>4</sup>Department of Theoretical Condensed Matter Physics, Universidad Autónoma de Madrid,  
12 E-28049 Madrid, Spain

13 <sup>5</sup>Russian Quantum Center, Skolkovo 143025, Russia

14 <sup>6</sup>Department of Physics, Moscow State University, 119991 Moscow, Russia

15 <sup>7</sup>Université de Bordeaux -CNRS, CELIA, UMR5107, F33405 Talence, France

16 <sup>8</sup>Technische Universität Berlin, Ernst-Ruska-Gebäude, Hardenbergstr. 36A, D-10623 Berlin,  
17 Germany

18 <sup>9</sup>Blackett Laboratory, Imperial College London, South Kensington Campus, SW7 2AZ London,  
19 United Kingdom

20 <sup>10</sup>Department of Physics, Humboldt University, Newtonstrasse 15, 12489 Berlin, Germany

22 **Strong field driven electric currents in condensed matter systems open**  
23 **new frontiers in petahertz electronics. In this regime new challenges arise**  
24 **as the role of the band structure and the quantum nature of electron-hole**  
25 **dynamics have yet to be resolved. Here we reveal the underlying attosec-**  
26 **ond dynamics that dictates the temporal evolution of carriers in multi-band**  
27 **solid state systems, via high harmonic generation (HHG) spectroscopy. We**  
28 **demonstrate that when the electron-hole relative velocity approaches zero,**  
29 **enhanced quantum interference leads to the appearance of spectral caustics**  
30 **in the HHG spectrum. Introducing the role of the dynamical joint density**  
31 **of states (JDOS) we identify its direct mapping into the spectrum, exhibit-**  
32 **ing singularities at the spectral caustics. By probing these singularities, we**  
33 **visualize the structure of multiple unpopulated high conduction bands. Our**  
34 **results open a new path in the control and study of attosecond quasi-particle**  
35 **interactions within the field dressed band structure of crystals.**

36 Induced by the strong field interaction, HHG provides a unique spectroscopic  
37 scheme to visualize the coherent evolution of petahertz currents inside solids.  
38 Since the first observation [1], solid HHG opened a door into the study of the  
39 electronic structure and dynamics in crystals [2, 3, 4, 5, 6, 7], multiple band dy-  
40 namics [8, 9, 10, 11] and complex many-body phenomena [12] in crystalline and  
41 amorphous systems [9]. For a moderate field strength the electron-hole dynamics  
42 are often described semi-classically by a single valence and conduction band of

---

\*These authors contributed equally to this work

43 the crystal. As we approach the strong field regime, new fundamental questions  
44 arise. What is the role of the band structure in such intense, ultrafast processes?  
45 How will electrons and holes interact on extremely short time scales, when they  
46 are still mutually quantum coherent? These questions pose some of the primary  
47 challenges in the emerging field of strong field interactions in solids.

48 In this paper we identify the quantum nature of the electron-hole wave-packet  
49 in solids, probing its strong-field attosecond dynamics over multiple bands. We  
50 observe enhanced quantum interference in the vicinity of Van Hove singularities  
51 [13, 14] and resolve the direct link between the dynamical JDOS and the HHG  
52 spectrum. Our study applies HHG spectroscopy in MgO, induced by a  $\lambda = 1.3\mu m$   
53 driving laser field. Adding a weak perturbative second harmonic (SH) field mod-  
54 ulates the internal dynamics, in a close analogy to a lock-in measurement. This  
55 scheme allows us to isolate extremely weak signals and identify the contribution  
56 of multiple band excitations, covering a spectral range of up to 30 eV. Our results  
57 show unequivocally that the internal dynamics related to HHG are dominated by  
58 the interband emission [1, 3, 4, 7, 15, 16, 17, 18, 19, 20, 21], which remains the  
59 dominant mechanism even when higher conduction bands are involved. We iden-  
60 tify the mapping between the dynamical JDOS at unique regions in the Brillouin  
61 zone (BZ) and the HHG spectrum. At Van Hove singularities, spectral caustics  
62 are induced, leading to a strong enhancement of the HHG signal. At these criti-  
63 cal points, the semi-classical picture fails [22], imprinting the dynamical quantum  
64 nature of the strong field interaction on the HHG spectrum.

65 The interband HHG mechanism can be viewed as a generalized electron-hole  
66 recollision process [15, 19], described by a semi-classical analysis [22]. Around

67 the peak of the laser field, an electron tunnels from the valence to the conduction  
68 band, forming an electron-hole pair. The laser field subsequently accelerates the  
69 pair, leading to their recombination and the emission of XUV radiation. This  
70 recollision model maps the semi-classical electron-hole trajectories into harmonic  
71 energies. The semi-classical analysis is not strictly limited to one conduction band  
72 – the strong laser field may excite the electron into higher bands, leading to the  
73 generation of complex spectral features in the HHG spectrum [8, 9, 10].

74 A fundamental aspect of strong-field induced tunneling is the localization of  
75 the excited electron-hole wave-packet in the BZ around the minimum band gap.  
76 As the wave-packet evolves under the influence of the laser field, it dynamically  
77 probes a narrow stripe of the BZ along the field’s polarization, as illustrated in  
78 figure 1e. Such localization is a key property of HHG spectroscopy – enabling the  
79 visualization of the band structure with unique angular resolution.

80 Resolving the angular dependence of the HHG spectrum is extremely chal-  
81 lenging – the HHG signal drops rapidly when the polarization is rotated off the  
82 main axes of the MgO crystal [23]. Such suppression becomes even more sig-  
83 nificant when the harmonics are produced from higher conduction bands [24]. In  
84 order to fully reveal the electron-hole dynamics, both its angular dependence as  
85 well as the contribution of multiple bands, we introduce an advanced measure-  
86 ment scheme that enables us to resolve and isolate the weak HHG signals.

87 Enhancing the detectability of weak signals by subjecting them to a known  
88 temporal modulation is a common practice in a wide range of applications, also  
89 known as lock-in measurement. We induce such a modulation by adding a weak  
90 SH field polarized perpendicular to the fundamental field’s polarization. So far

91 this scheme has been applied to probe the internal dynamics in gas phase HHG  
92 [25] (and references within) and interband currents in ZnO [15]. Scanning the  
93 two-color delay leads to a periodic modulation of the HHG spectrum with four  
94 times the fundamental laser's frequency. Fourier analysis isolates the oscillating  
95 component, resolving the extremely weak HHG signals, buried in a large exper-  
96 imental background. In addition, XUV emission from higher conduction bands  
97 can be highly susceptible to the two-color delay [26], resulting in an enhanced  
98 signature in the oscillating component. Figures 1a, 1b and 1c compare the Fourier  
99 amplitudes of the modulated HHG signals and the spectra averaged over the two  
100 color delay for crystal orientations of  $0^\circ$ ,  $33^\circ$  and  $45^\circ$ , respectively (see figure 1d).  
101 Clearly, the averaged spectrum is accompanied by large background noise and is  
102 dominated by harmonic 17 ( $\sim 16.2$  eV) for  $33^\circ$  and  $45^\circ$  and harmonics 17 and 19  
103 ( $\sim 18.1$  eV) for  $0^\circ$ . Resolving the oscillating spectrum by Fourier analysis shows a  
104 dramatic enhancement of the signal over the noise. This measurement exhibits the  
105 appearance of new spectral components, which were so far hidden in the averaged  
106 spectrum.

107 The oscillating spectra reveal that for all crystal angles, the spectrum extends  
108 beyond the first band edge ( $\sim 18$  eV), indicating the strong contribution of higher  
109 conduction bands. The most important observation is associated with the angular  
110 dependence of the measurement – the oscillating spectrum varies drastically and  
111 surprisingly for different angles exhibiting enhanced features and structures over  
112 a wide spectral range. We identify three distinct structures: enhanced harmonics  
113 around 16-17 eV extending to 18 eV at  $0^\circ$ , an increased signal in the 20.5-22.5 eV  
114 region at  $33^\circ$  and a clear spectral feature at 23.5-26 eV, narrowing at  $45^\circ$ . These

115 features can not be associated with the spectral response of the dipole coupling  
 116 between the bands, due to their moderate variation with energy (see SI). What  
 117 is the origin of the enhancement mechanism in these spectral regions and what  
 118 information does it provide?

119 We start by analyzing the enhancement at 23.5-26 eV. This observation is strik-  
 120 ing – clearly, this spectral region originates from a high conduction band excitation  
 121 associated with a low population transfer. Furthermore, while the oscillating spec-  
 122 trum changes dramatically with the crystal orientation, this feature remains robust.  
 123 Such observation suggests that it originates from areas of the BZ which are orien-  
 124 tation independent. These areas can be found around the  $\Gamma$  point of the BZ which  
 125 is common to all crystal orientations. Figures 2a and 2b describe the band struc-  
 126 ture for crystal orientations of  $0^\circ$  and  $45^\circ$ , respectively. Indeed, we find two high  
 127 conduction bands intersecting the  $\Gamma$  point at an energy of 24.57 eV. At the  $\Gamma$  point  
 128 these bands, as well as any band, have a critical point,  $\nabla_{\mathbf{k}}E_c(\mathbf{k}) = \nabla_{\mathbf{k}}E_v(\mathbf{k}) = 0$ ,  
 129 such that the gradient of the energy gap is zero as well:

$$\nabla_{\mathbf{k}}(E_c(\mathbf{k}) - E_v(\mathbf{k})) = \nabla_{\mathbf{k}}\varepsilon_g(\mathbf{k}) = 0 \quad (1)$$

130 where  $\mathbf{k}$  is the lattice momentum,  $E_c$  is the conduction band and  $E_v$  is the va-  
 131 lence band. At the critical points, where  $\nabla_{\mathbf{k}}\varepsilon_g(\mathbf{k}) = 0$ , the JDOS of a crystalline  
 132 solid becomes singular. These singularities are known as Van Hove singulari-  
 133 ties and most commonly arise in the analysis of optical absorption and reflection  
 134 spectra [27, 28]. In contrast to these measurements, performed over long time  
 135 scales, HHG in bulk crystal is attributed to the sub-cycle electronic dynamics  
 136 where the electron and hole wave-packets remain mutually coherent. In addition,

137 these wave-packets, which are initiated by tunneling, are localized in the BZ. As  
 138 a result, the harmonic emission can not be understood by merely counting the  
 139 total number of available states throughout the BZ. Instead, the quantum nature  
 140 of the singularity has to be considered at specific  $\mathbf{k}$  regions. During such short  
 141 time scales, we can view the singularities as points where the relative velocity of  
 142 the electron and hole wave-packets is zero. This results in enhanced interference  
 143 effects, encoding dynamical quantum information into the HHG spectrum.

144 The quantum expression for the interband currents can be written as [19, 22]

$$\mathbf{j}_{er}(\omega) = \omega \int_{B.Z.} d^3\mathbf{k} \iint \mathbf{g}(\mathbf{k}, t', t) \exp^{-iS(\mathbf{k}, t', t) + i\omega t} dt dt' \quad (2)$$

$$S(\mathbf{k}, t', t) = \int_{t'}^t \varepsilon_g(\mathbf{k} - \mathbf{A}(t) + \mathbf{A}(\tau)) d\tau \quad (3)$$

145 where  $\mathbf{A}$  is the vector potential and  $\omega$  the harmonic energy.  $g(\mathbf{k}, t', t)$  represents  
 146 the slowly varying term compared to the fast oscillating exponential containing  
 147 the semi-classical action,  $S(\mathbf{k}, t', t)$ . Under these conditions we can use the sta-  
 148 tionary phase approximation (SPA) for  $\mathbf{j}_{er}(\omega)$  which defines the semi-classical  
 149 mapping between harmonic energy and recolliding electron and hole trajectories.  
 150 The interband current for a single stationary solution as obtained from the SPA, is  
 151 proportional to  $\sqrt{|\hat{\mathbf{S}}''(\mathbf{k}_{st}, t'_{st}, t_{st})|^{-1}}$ , where  $\hat{\mathbf{S}}''(\mathbf{k}_{st}, t'_{st}, t_{st})$  is the Hessian matrix  
 152 of the function  $S(\mathbf{k}, t', t)$  at the stationary points. This derivation directly links the  
 153 HHG spectrum and the band structure (for a detailed derivation see SI):

$$I(\omega) \propto |\mathbf{j}_{er}(\omega)|^2 \propto \omega^2 \left| \sum_{\mathbf{k}_{st}} \frac{\tilde{\mathbf{g}}(\mathbf{k}_{st}, t'_{st}, t_{st}) e^{-iS(\mathbf{k}_{st}, t'_{st}, t_{st})}}{\sqrt{|\nabla_{\mathbf{k}} \varepsilon_g(\mathbf{k}_{st})|}} \right|^2 \quad (4)$$

154 where  $\varepsilon_g(\mathbf{k}_{st})$  is the energy difference between electron and hole at the time of rec-  
 155 ollision, which defines the harmonic energy, and  $\tilde{\mathbf{g}}$  accounts for all pre-exponential  
 156 terms. The expression for the spectral intensity strongly resembles that of the  
 157 JDOS, except it is weighted by the exponent of the semi-classical action, there-  
 158 fore, it is associated with a *dynamical JDOS*. This exponent gives the quantum  
 159 phase and amplitude associated with each trajectory. Due to strong field tunnel-  
 160 ing, the quantum amplitude strongly attenuates with increasing  $|\mathbf{k}_{st}^\perp|$ ,  $\mathbf{k}_{st}^\perp$  being  
 161 the component of  $\mathbf{k}_{st}$  perpendicular to the laser's polarization. Such attenuation  
 162 expresses the fact that HHG indeed originates from a narrow stripe of the BZ, pro-  
 163 viding the significant angular dependence of the spectrum. Clearly this mapping  
 164 becomes singular at the extrema of the energy gap as described by equation (1).

165 The singularity in the mapping between harmonic energy and electron-hole  
 166 trajectories can be described within the framework of caustics [29]. Caustics  
 167 are universal phenomena in nature that link processes observed in many differ-  
 168 ent branches of physics. Previous studies identified the appearance of caustics in  
 169 gas phase HHG [29, 30], where they reveal the quantum nature of the process in  
 170 a regime where classical analysis fails. Figure 1e illustrates the origin of spec-  
 171 tral caustics in condensed matter systems, when the electron-hole wave-packet  
 172 has zero relative velocity at the extrema of the band gap. We find that spectral  
 173 caustics reveal the rich, quantum, spatio-temporal nature of electronic dynamics  
 174 in solids, dictated by the direct link between the band structure and strong field  
 175 interaction.

176 The enhancement at 23.5-26 eV can be identified as a spectral caustic orig-  
 177 inating from the  $\Gamma$  point. Figures 2a, 2d present the band structure for crystal

178 orientations of  $0^\circ$  and  $45^\circ$ , respectively. Another energy band intersecting the  $\Gamma$   
 179 point can be found at 18.8 eV (figure 2d), leading to a robust spectral feature near  
 180 18 eV as can be seen in figures 1a-c. While the crossing of energy bands at the  
 181  $\Gamma$  point will always result in singularities, spectral caustics can be found at other  
 182 points in the BZ as well. Along the high symmetry axes of the crystal,  $0^\circ$  and  
 183  $45^\circ$ , each band-gap consists of several critical points where  $\nabla_{\mathbf{k}}\varepsilon_g(\mathbf{k}_{st}) = 0$ . In  
 184 figure 2 we show how these points are directly imprinted in the experimentally  
 185 resolved HHG spectrum. Figures 2b, 2e describe the gradient of each band gap as  
 186 a function of the band gap energy, for crystal orientations of  $0^\circ$  and  $45^\circ$ , respec-  
 187 tively. Figures 2c, 2f present the measured oscillating spectra at these angles. The  
 188 highlighted lines in figures 2c, 2f emphasize the energy points where the gradi-  
 189 ent is zero and the dynamical JDOS becomes singular according to equation (4) .  
 190 Indeed, caustics dominate the brightest features in the HHG spectrum, leading to  
 191 a dramatic spectral focusing and enhancement of the weak signal associated with  
 192 higher conduction bands excitation.

193 While the singularities are observed along the high symmetry axes of the crys-  
 194 tal, figure 3a shows that enhanced spectral features are observed at other angles  
 195 as well . Their origin can be understood through the dynamical JDOS, by looking  
 196 at the gradient of the different band gaps,  $\nabla_{\mathbf{k}}\varepsilon_g(\mathbf{k})$ . Figures 3b and 3c present  
 197 2D images of the gradients of bands 5 and 7. In both bands we can identify a  
 198 pronounced valley at specific  $\mathbf{k}$  values and across a large angular range. Along  
 199 these valleys  $\nabla_{\mathbf{k}}\varepsilon_g(\mathbf{k})$  is small, leading, according to equation (4) to a local spec-  
 200 tral enhancement in the HHG signal. Specifically, the gradient valley at band 5  
 201 is mapped to a pronounced spectral enhancement around 16-17 eV, while the val-

202 ley in band 7 leads to the enhanced spectral feature in the 20.5-22.5 eV region,  
203 marked by the violet and cyan areas, respectively. Importantly, their observation  
204 is robust with the crystal orientation, mapping the angular variation of the gradient  
205 valleys.

206 Understanding the mapping between the band structure and the HHG spectrum  
207 allows us to study the fundamental dynamical properties of strong field dressed  
208 phenomena in solids [11, 31]. Such effects play an important role when the band  
209 structure becomes degenerate. In the vicinity of these regions the intricate inter-  
210 play between the strong field dressing and the orientation dependent band cou-  
211 plings can have a dramatic effect on the electronic dynamics. At the  $\Gamma$  point of  
212 MgO there is a two band degeneracy at 24.6 eV (bands 9 and 10, plotted in green  
213 and in red), protected by the crystal symmetries. Taking a closer look at the asso-  
214 ciated caustic we find that our measurement encodes a signature of such a strong  
215 field mechanism. The enhancement observed at the high energy end of the spec-  
216 trum, 23.5-26 eV, is robust with crystal orientation, as one would expect from a  
217 feature originating from the  $\Gamma$  point. However, the spectral shape of this feature  
218 shows a striking angular dependence, where the peak shifts from harmonic 25 at  
219  $0^\circ$  to 26 at  $45^\circ$ , exhibiting complex spectral structures at intermediate angles (see  
220 dashed line in figure 3a). By rotating the crystal we change the couplings and  
221 field dressing of these two bands, imprinting the intricate underlying strong field  
222 dynamics into the angular dependence of the HHG spectrum.

223 Our results establish the fundamental connection between the electronic struc-  
224 ture of the crystal and the strong field process. This study reveals how strong  
225 field attosecond metrology serves as an extremely sensitive probe of ultrafast dy-

226 namical quantum interference between electron-hole wave-packets in solids. We  
227 identify the important role of the dynamical JDOS, dictated by the strong field  
228 nature of the interaction. As the dynamical JDOS becomes singular, the trajec-  
229 tory picture that governs the interband HHG process fails, giving rise to spectral  
230 caustics. These findings provide a unique insight into the dressing of the band  
231 structure by the strong laser field. The mapping between the HHG spectrum and  
232 the band gaps serves as unequivocal evidence to the dominant role of interband  
233 emission over numerous conduction bands. Looking forward, our study will form  
234 the framework for a large range of attosecond scale phenomena, opening a new  
235 path in the study of quasi-particle interactions within the field dressed band struc-  
236 ture of crystals. It will allow the study of ultrafast dynamics such as electron-hole  
237 interactions, leading to the formation of excitons, or electron-electron-phonon in-  
238 teractions. Furthermore, the control over sub-cycle electronic currents will play  
239 a role in the establishment of compact solid state XUV sources, as well as in the  
240 field of petahertz electronics.

## 241 **References**

- 242 [1] Ghimire, S. *et al.* Observation of high-order harmonic generation in a bulk  
243 crystal. *Nature physics* **7**, 138 (2011).
- 244 [2] Vampa, G. *et al.* All-optical reconstruction of crystal band structure. *Physi-  
245 cal review letters* **115**, 193603 (2015).

- 246 [3] Luu, T. T. *et al.* Extreme ultraviolet high-harmonic spectroscopy of solids.  
247 *Nature* **521**, 498 (2015).
- 248 [4] Garg, M. *et al.* Multi-petahertz electronic metrology. *Nature* **538**, 359  
249 (2016).
- 250 [5] Liu, H. *et al.* High-harmonic generation from an atomically thin semicon-  
251 ductor. *Nature Physics* **13**, 262 (2017).
- 252 [6] Yoshikawa, N., Tamaya, T. & Tanaka, K. High-harmonic generation in  
253 graphene enhanced by elliptically polarized light excitation. *Science* **356**,  
254 736–738 (2017).
- 255 [7] Hohenleutner, M. *et al.* Real-time observation of interfering crystal electrons  
256 in high-harmonic generation. *Nature* **523**, 572 (2015).
- 257 [8] Ndabashimiye, G. *et al.* Solid-state harmonics beyond the atomic limit. *Na-*  
258 *ture* **534**, 520 (2016).
- 259 [9] You, Y. S. *et al.* High-harmonic generation in amorphous solids. *Nature*  
260 *communications* **8**, 724 (2017).
- 261 [10] You, Y. S. *et al.* Laser waveform control of extreme ultraviolet high harmon-  
262 ics from solids. *Optics letters* **42**, 1816–1819 (2017).
- 263 [11] Hawkins, P. G., Ivanov, M. Y. & Yakovlev, V. S. Effect of multiple conduc-  
264 tion bands on high-harmonic emission from dielectrics. *Physical Review A*  
265 **91**, 013405 (2015).

- 266 [12] Silva, R., Blinov, I. V., Rubtsov, A. N., Smirnova, O. & Ivanov, M. High-  
267 harmonic spectroscopy of ultrafast many-body dynamics in strongly corre-  
268 lated systems. *Nature Photonics* 1 (2018).
- 269 [13] Ashcroft, N. W. & Mermin, N. D. Solid state physics (holt, rinehart and  
270 winston, new york, 1976) 144–145.
- 271 [14] Van Hove, L. The occurrence of singularities in the elastic frequency distri-  
272 bution of a crystal. *Physical Review* **89**, 1189 (1953).
- 273 [15] Vampa, G. *et al.* Linking high harmonics from gases and solids. *Nature* **522**,  
274 462 (2015).
- 275 [16] Schubert, O. *et al.* Sub-cycle control of terahertz high-harmonic generation  
276 by dynamical bloch oscillations. *Nature Photonics* **8**, 119 (2014).
- 277 [17] Kemper, A., Moritz, B., Freericks, J. & Devereaux, T. Theoretical descrip-  
278 tion of high-order harmonic generation in solids. *New Journal of Physics* **15**,  
279 023003 (2013).
- 280 [18] Higuchi, T., Stockman, M. I. & Hommelhoff, P. Strong-field perspective  
281 on high-harmonic radiation from bulk solids. *Physical review letters* **113**,  
282 213901 (2014).
- 283 [19] Vampa, G. *et al.* Theoretical analysis of high-harmonic generation in solids.  
284 *Physical review letters* **113**, 073901 (2014).

- 285 [20] Golde, D., Meier, T. & Koch, S. High harmonics generated in semiconduc-  
286 tor nanostructures by the coupled dynamics of optical inter-and intraband  
287 excitations. *Physical Review B* **77**, 075330 (2008).
- 288 [21] Tancogne-Dejean, N., Mücke, O. D., Kärtner, F. X. & Rubio, A. Impact of  
289 the electronic band structure in high-harmonic generation spectra of solids.  
290 *Physical review letters* **118**, 087403 (2017).
- 291 [22] Vampa, G., McDonald, C., Orlando, G., Corkum, P. & Brabec, T. Semiclas-  
292 sical analysis of high harmonic generation in bulk crystals. *Physical Review*  
293 *B* **91**, 064302 (2015).
- 294 [23] You, Y. S., Reis, D. A. & Ghimire, S. Anisotropic high-harmonic generation  
295 in bulk crystals. *Nature Physics* **13**, 345 (2017).
- 296 [24] Wu, M. *et al.* Orientation dependence of temporal and spectral properties of  
297 high-order harmonics in solids. *Physical Review A* **96**, 063412 (2017).
- 298 [25] Bruner, B. D. *et al.* Multidimensional high harmonic spectroscopy. *Journal*  
299 *of Physics B: Atomic, Molecular and Optical Physics* **48**, 174006 (2015).
- 300 [26] Li, J.-B. *et al.* Enhancement of the second plateau in solid high-order har-  
301 monic spectra by the two-color fields. *Optics Express* **25**, 18603–18613  
302 (2017).
- 303 [27] Ehrenreich, H. & Philipp, H. Optical properties of ag and cu. *Physical*  
304 *Review* **128**, 1622 (1962).

- 305 [28] Roessler, D. & Walker, W. Electronic spectrum and ultraviolet optical prop-  
306 erties of crystalline mgo. *Physical Review* **159**, 733 (1967).
- 307 [29] Raz, O., Pedatzur, O., Bruner, B. D. & Dudovich, N. Spectral caustics in  
308 attosecond science. *Nature Photonics* **6**, 170 (2012).
- 309 [30] Faccialà, D. *et al.* Probe of multielectron dynamics in xenon by caustics in  
310 high-order harmonic generation. *Physical review letters* **117**, 093902 (2016).
- 311 [31] McDonald, C., Vampa, G., Corkum, P. & Brabec, T. Intense-laser solid state  
312 physics: unraveling the difference between semiconductors and dielectrics.  
313 *Physical review letters* **118**, 173601 (2017).

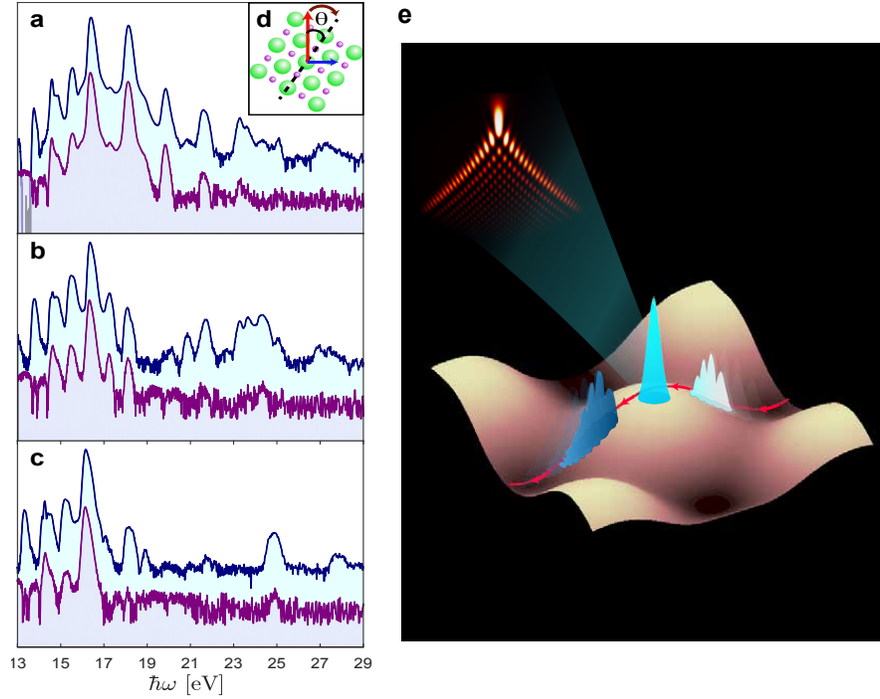


Figure 1: **HHG spectroscopy in MgO.** **a, b, c** normalized oscillating HHG spectrum (cyan) and the normalized averaged spectrum (violet) in logarithmic scale for orientations **a**  $0^\circ$ , **b**  $33^\circ$  and **c**  $45^\circ$ . The two spectra are vertically shifted to allow better visibility. **d** the crystal is oriented to angle  $\theta$  with respect to the fundamental field's polarization (red arrow) while the second harmonic (blue arrow) is orthogonally polarized. **e** Schematic description of enhanced quantum interference, leading to spectral caustics. The white, cyan and blue wave-packets are snapshots of the combined electron-hole wave-function **at three different times**, probing the band gap of band 10 (figure 2d) along the laser polarization (red line). The modulated white and blue snapshots capture the wave-packet where the electron-hole relative velocity is non-zero. At the extremum of the gap, where the relative velocity is zero (cyan snapshot), enhanced interference leads to the emission of a bright spectral caustic (illustrated by the red interference pattern).

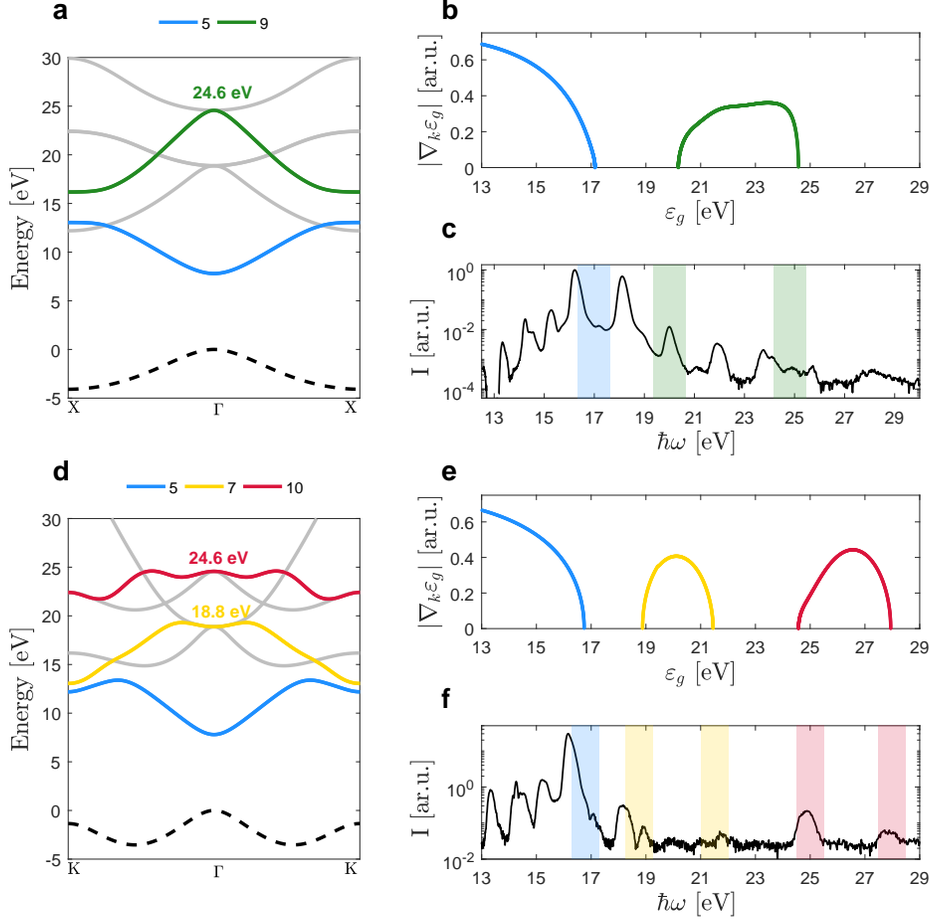


Figure 2: **Spectral caustics at singular points of the dynamical JDOS.** The band structure along the  $0^\circ$  ( $\Gamma - X$ ) (a) and  $45^\circ$  ( $\Gamma - K$ ) (d) orientations, respectively. The dashed line is the valence band and the colored lines are the conduction bands which are dipole coupled to the valence band. The gray bands indicate the existing bands which are not dipole coupled. b and e plot  $|\nabla_{\mathbf{k}} \varepsilon_g(\mathbf{k})|$  along  $0^\circ$  and  $45^\circ$ , respectively. The energies for which  $|\nabla_{\mathbf{k}} \varepsilon_g(\mathbf{k})| = 0$ , where singularities in the dynamical JDOS occur, are highlighted by corresponding colored bars on the oscillating spectra c ( $0^\circ$ ) and f ( $45^\circ$ ), emphasizing their link to the spectral caustics.

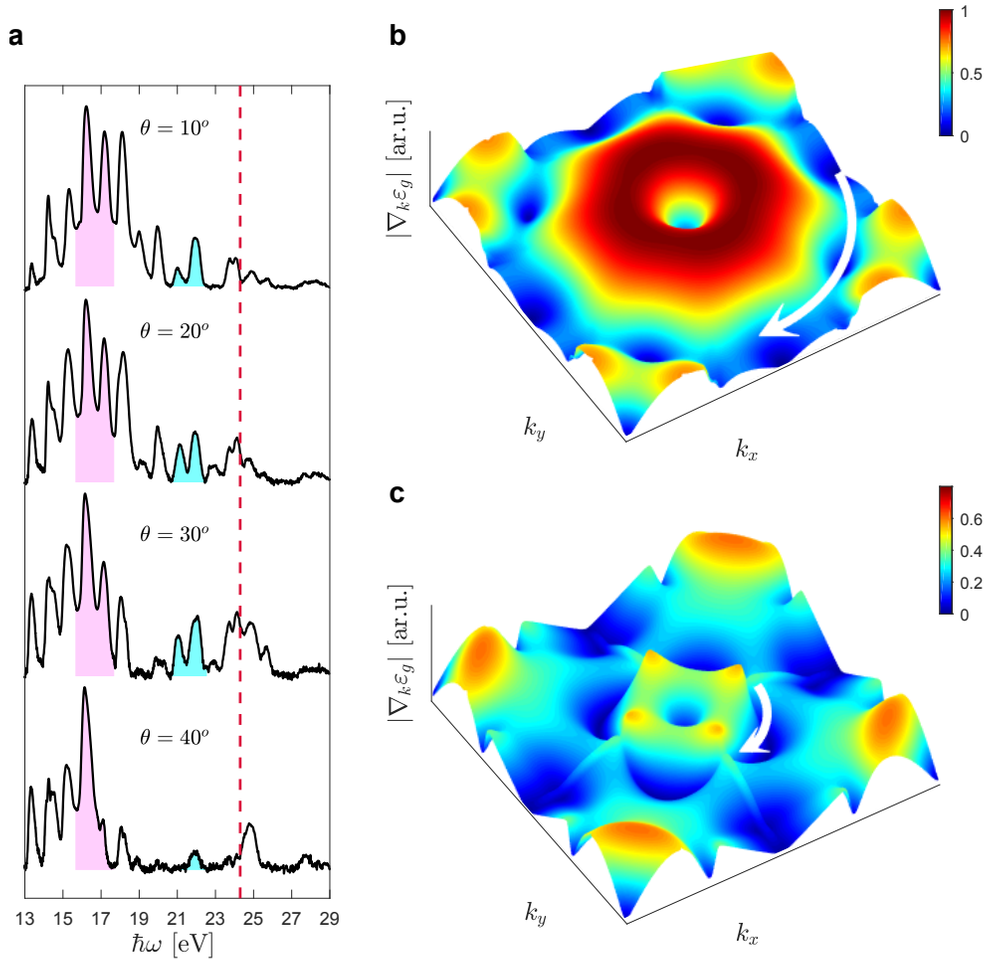


Figure 3: **HHG spectral enhancement at high dynamical JDOS valleys.** **a** the oscillating spectrum for  $\theta = 10^\circ, 20^\circ, 30^\circ, 40^\circ$ . Two enhanced spectral structures which persist over many orientations are marked by violet and cyan. These spectral structures originate from high dynamical JDOS. The dashed red line marks the spectral caustic between 23.5-26 eV highlighting its angular dependence. **b** and **c** are the 2D gradients of the band gap for bands 5 and 7, respectively. The arrows show valleys of low  $\nabla_{\mathbf{k}}\varepsilon_g(\mathbf{k})$ , i.e. high dynamical JDOS, corresponding to the enhanced spectral features in **a**. The valley in **b** corresponds to the structure marked by violet and **c** to the one in cyan.