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Can a shock-induced phonon up-pumping model relate to impact sensitivity of molecular crystals, polymorphs and cocrystals?†

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Impact sensitivity engineering of high-energy molecular crystals requires accurate predictive models. For this purpose, the promising multi-phonon based approach is selected, assessing a bit more its strengths and weaknesses. Presently used with high-quality phonon calculations of 22 molecular crystals, using a physics-based criterion to determine the phonon bath extent, the resulting intrinsic shock sensitivity index (SSI) is compared to the most common marker of impact sensitivity, h_{50} , as determined from drop-weight impact tests. Selecting a data subset from experiments performed under very similar conditions (2.5 kg hammer with grit and 30–40 mg samples), the model can predict h_{50} values for mono-molecular crystals with very good accuracy, including the ability to discriminate the polymorphs of HMX and CL20. This very good agreement validates an initial indirect up-pumping mechanism occurring under these conditions, where the doorway modes also interact with the phonon bath. However, the phonon bath criterion for mono-molecular crystals does not transfer well to cocrystals. Owing to the vibrational coupling of the co-molecules, it seems a broader phonon bath should be considered. Additionally recalling experimental uncertainty and various experimental factors affecting h_{50} values for a given compounds, we recommend that the density of the sample, granularity and morphology be systematically considered and reported along with measurements, which will in turn allow for more systematic data and predictive capabilities for sensitivity models.

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I. Introduction

Tailoring the impact sensitivity of high-energy molecular crystals is important for developing a quantitative perspective on use and safety protocols across different applications. In general, an ideal balance between maximum power¹ and acceptable sensitivity^{2–4} is hard to achieve without a multiscale approach to control key molecular and mesoscale drivers. Controlling the explosive sensitivity can be achieved in many ways, including using nanograins,^{5–10} a coating of micrograins^{11–13} for desensitization, or changing the molecular blocks and their packing using co-crystallization.^{2,14–18} Currently, a common route to create cocrystals is through an experimental trial-and-error process. Then, further experiments can be performed to rank their sensitivity with respect to their

pure compounds. But with the increasing trend of computational design,¹⁹ it is tempting to predict such cocrystals alongside their full properties. The thermomechanical properties and shock properties can be well-predicted from *ab initio* methods using energy and dispersion corrected DFT-D,²⁰ but predicting shock sensitivity remains challenging.

The most common marker of impact sensitivity is the height, denoted h_{50} , at which a dropped weight ignites a sample with 50% probability (or the related energy E_{50}). The large database of experimental h_{50} , covering hundreds of molecular crystals, has helped in devising many data-centric models. These models give insights about some trends with various factors pertaining to molecular formula or to physical properties of the crystal. But scarce are the ones outperforming 80% accuracy. A few most recent models are shortly described hereafter. Jensen *et al.*²¹ have calculated bond dissociation energy, heat and temperature of detonation of 70 isolated, nitroaromatic molecules to come up with a model with a correlation coefficient R^2 between 0.81 and 0.85. Mathieu *et al.*^{22,23} have retrieved numerous data for 308 energetic molecules to devise a model based on both thermodynamic and kinetic descriptors, with a correlation coefficient R^2 of 0.8. Xiong *et al.*²⁴ have a similar but less accurate model, incorporating the self-sustaining ignition ability of 150 energetic molecules, and reaching a Pearson correlation coefficient of 0.67 (meaning an R^2 value around 0.45). Nevertheless,

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† Electronic supplementary information (ESI) available: (1) Evaluation of the “nitroaromatics” model for nitramines; (2) phonon density of states of β -HMX from DFT-D – issues when using only the Gamma point; (3) optimized lattice parameters from DFT-D. vDOS of the molecular crystals and of the extracted molecules in gas phase, and some mode frequency assignments. Zip file of the vDOS data for the molecular crystals. See DOI: <https://doi.org/10.1039/d2ra05062e>



Mathieu *et al.* hypothesize in their papers many sources of discrepancy from experimental h_{50} , and propose to add a crystal packing coefficient for a possible improvement. In an attempt to go beyond mere molecular properties, Bondarchuk²⁵ has considered solid-state properties of 24 energetic crystals, calculated the pressure at which the electronic band gap is lower than 1 meV, and ended up with a shock sensitivity model with a correlation coefficient R^2 of 0.83. The method is a bit expensive, and considering the bulk modulus instead may be more efficient, as done by Deng *et al.*²⁶ for 240 nitroaromatics. They achieve a correlation coefficient R^2 of 0.91. However, our application of their model to nitramines – see Section 1 in our ESI – is acceptable only for nitroaromatics, like TATB and TNT, but deeply erroneous for the non-aromatic nitramines: for instance, β -HMX is predicted with an h_{50} higher than 800 cm, which is much less sensitive than even TATB. Considering polymorphs, this model erroneously predicts β - and δ -HMX similarly insensitive, whereas both are actually sensitive, with δ -HMX even more so.²⁷ Then, β - and ε -CL20 are predicted with sensitivity similar to TNT, whereas both are actually more sensitive than that, and β -CL20 more than ε -CL20.²⁸ This model lacks transferability and it would need a parameterization for nitramines. Alternatively, extending it to a broader range of energetic crystals than initially intended will likely result in a decreased correlation coefficient R^2 , probably around 0.8, like the other models.

Using vibrational properties of the molecular crystals, an approach based on phonon interactions and up-pumping,^{29–31} and improved by Michalchuk *et al.*^{32,33} looks promising. In their studies, they recall the theoretical approach to up-pumping with the following assumptions. The shock energy is first transferred to the phonon bath, the low frequency modes within the range $[0, \omega_{\max}]$ related to intermolecular or lattice vibrations and some amalgamated intramolecular modes. Then, this energy is transferred to the “doorway” modes, whose frequencies are within the range $[\omega_{\max}, 2\omega_{\max}]$, and they can be subsequently up-pumped to higher frequencies related to intramolecular vibrations until bond rupture. In their model, ω_{\max} depends on the distribution of low-frequency modes in a given material, generally around 200 cm^{−1}. A few approximations, such as *ab initio* calculation of the vibrational density of states (vDOS) restricted to the Gamma point and, from it, the derivation of multiphonon spectra, make this method affordable. Overall, their model was applied to less than 20 energetic materials and the sensitivity prediction aligns well with experimental h_{50} , albeit without explicitly quantifying a correlation coefficient. Their model succeeds in ranking δ -HMX as more sensitive than β -HMX.³⁴ *Ab initio* calculations restricted to the Gamma point can be motivated by high throughput considerations, but this approximation may not be suitable to discriminate other molecular crystals, co-crystals and their polymorphs. Indeed, we show in Section 2 of the ESI† that the vDOS of β -HMX (Fig. S1†) obtained using this approximation strongly differs from the ones using a larger number of k-points or supercells, and especially in the low-frequency range. A good description of this range associated with lattice/intermolecular vibrations is crucial for molecular crystals,³⁵ both in frequencies

and relative amplitudes. It has been shown that paying attention to this critical range can resolve large discrepancies between theory and experiments.²⁰

Drott and Fayer³⁰ posited that a vibrational up-pumping scheme describing the internal energy flow in shocked organic materials occurs within a few tens of picoseconds after the shock front. This approach cannot *a priori* relate to impact experiments, where the deformations or compressions do not always generate a shockwave. The shock-less impact is thus commonly accepted. Storm *et al.*³⁶ recall the large differences in time scale and pressure between shock and impact experiments (0.05–2 μ s and 3–20 GPa *versus* 200–250 μ s and 0.7–1.5 GPa). The thermalization of the phonon bath is extremely fast, around 1 ps (ref. 37) or less,³⁸ which is significantly faster than internal-mode lifetime of a few nanoseconds.^{37,38} Analyzing 21 explosives, their h_{50} from impact experiments and their P_{90} (pressure to initiate an explosive pressed to 90% of its theoretical maximum density) from shock experiments, Storm *et al.*³⁶ actually find an excellent correlation ($R^2 = 0.99$, excluding 3 outliers). This high correlation supports the approach for predicting impact sensitivity from shock-induced phonon up-pumping.

From the high-quality phonon spectra of 20 mono-molecular (molecules of one type) crystals, including some polymorphs and 2 cocrystals, this paper explores the capabilities of the multi-phonon based approach to predict their impact sensitivity from a shock sensitivity index (SSI), with a thorough comparison to experimental h_{50} and observations. One difficulty in the up-pumping scheme is the selection of the phonon bath extent ω_{\max} .³² The criterion of a frequency gap to define the upper limit of the phonon bath was found not fully satisfactory, and we presently use a more physics-based criterion. The model and its capabilities to discriminate polymorphs and cocrystals are discussed, acknowledging sources of discrepancy in theory and on the experimental side.

II. Methods

A. DFT-D and variable-cell relaxation

First principle calculations are carried out using the Density Functional Theory (DFT) as implemented in Quantum Espresso 6.7 (QE),³⁹ the Perdew–Burke–Ernzerhof (PBE) generalized gradient approximation (GGA) functional, the Grimme D2 correction for a proper description of the van der Waals interactions (denoted hereafter PBE-D2) and the ultrasoft plane-wave pseudopotential (PP) from PSLibrary 1.0.0 (ref. 40) with non-linear core correction (NLCC). The energy cutoff for wave functions is set at 90 Ry for all molecular crystals and the cutoff for charge density is set at 900 Ry. A convergence threshold of 10^{-8} Ry is used for Self-Consistent Field (SCF) calculations. The Brillouin zone is sampled using a Monkhorst–Pack off-grid k-point mesh adapted to every crystal and cocrystal reported in the ESI, Table S3.† Performing variable-cell relaxation at zero pressure, the convergence is monitored using the Broyden–Fletcher–Goldfarb–Shanno (BFGS) algorithm for both lattice parameters and atom positions, using thresholds of 10^{-5} Ry for the system energy, 10^{-4} Ry bohr^{−1} for the forces and 0.1 kbar for



the pressure tensor. The resulting lattice parameters, reported in Table S4 of the ESI,[†] are in excellent agreement with ambient experimental data. The ability of PBE-D2 and PP to reproduce ambient structures of explosive molecular crystals is known,^{20,41} as well as the suitability for accurate determination of their phonon density of state.²⁰

B. Vibrational density of state

Phonon properties are determined using atomic finite displacements and the supercell method as implemented in Phonopy 2.9.1,⁴² with supercells as specified in Table S3.[†] For every optimized structure, Phonopy generates supercell configurations where the atoms of a given vibrational mode are displaced by 0.02 bohr. Using the molecule symmetry and the group symmetry of each crystal (Table S4[†]), these modes are reduced to the number of irreducible representations reported in Table S3.[†] The force constants are then calculated by QE (a single off-grid k-point is enough, due to supercell replication), and post-processed by Phonopy to obtain the vDOS, also denoted $g(\omega)$, using a dense $16 \times 16 \times 16$ q-point mesh grid and a Gaussian broadening of 2 cm^{-1} . The vDOS bin size is 1 cm^{-1} , and vDOS values below a threshold of 10^{-4} states per cm^{-1} are considered zero. Note that, as expected, the vDOS integration correctly yields three times the number of atoms per unit cell ($3N$). Fig. S2 in the ESI[†] compares the vDOS of β -HMX, α -NTO, TATB and α -FOX-7 to experimental spectra of inelastic neutron scattering (INS) performed at 20 K.^{32,43} Peak positions are in very good agreement (see Section 4 of the ESI[†]). In addition, to visualize the phonon modes and to quickly assign them to their corresponding frequencies, vDOS at the Γ q-point are calculated for every molecular crystal and for the extracted molecules in gas phase. For the latter, the extracted molecules are not geometrically re-optimized, but kept in the conformation they had in the crystal, and placed in a box of at least twice the molecule size. An example for β -HMX is provided in Fig. 1(a). It can be seen that, as expected, the frequencies of the extracted

molecule in gas phase are lower than in the crystal, and especially for the phonon bath. Also, some molecular modes can degenerate in the crystal,⁴⁴ and some others can synchronize.

C. Sensitivity prediction from the overtone-based model

Following our effort to obtain accurate vDOS for molecular crystals, especially in the low-frequency range, it makes sense to use methods based on a multi-phonon approach to model their shock sensitivity. As described by Michalchuk *et al.*,³² the lattice phonon bath (frequency $\omega < \omega_{\max}$, where ω_{\max} depends on the given molecular crystal) is excited and stores the initial energy of the shock. This amount of energy is then rapidly up-pumped to intra-molecular modes, quicker than dissipation, and some of these vibrational modes are responsible for bond breaking. In the doorway mode approach, the initial stage of up-pumping is dominated by processes involving two “bath” phonons converting into one intra-molecular phonon (the doorway modes are thus defined by $\omega_{\max} < \omega < 2\omega_{\max}$). The excited doorway modes can subsequently interact with the phonon bath again, building an indirect thermal mechanism, and up-pumping is thus considered up to $3\omega_{\max}$.³³ Direct interactions between three “bath” phonons are also considered within the same limit of $3\omega_{\max}$.

Michalchuk *et al.*³² have proposed more sophisticated variants of their own method. This study focuses on the simplest version: the overtone (OT) based model. Its advantage is that it requires only one empirically adjusted parameter: the extent ω_{\max} of the phonon bath. In an attempt to minimize the human factor, ω_{\max} is adjusted the same way for all molecular crystals. From our observation of the calculated vDOS and the assigned vibrational modes, we have noticed that the best correlation and consistency across the board is achieved when ω_{\max} corresponds to the X-NO_2 twisting modes (see example in Fig. 1(a)). The present work thus focuses on this physics-based criterion applied to all the crystals hereafter. Note that a comparative

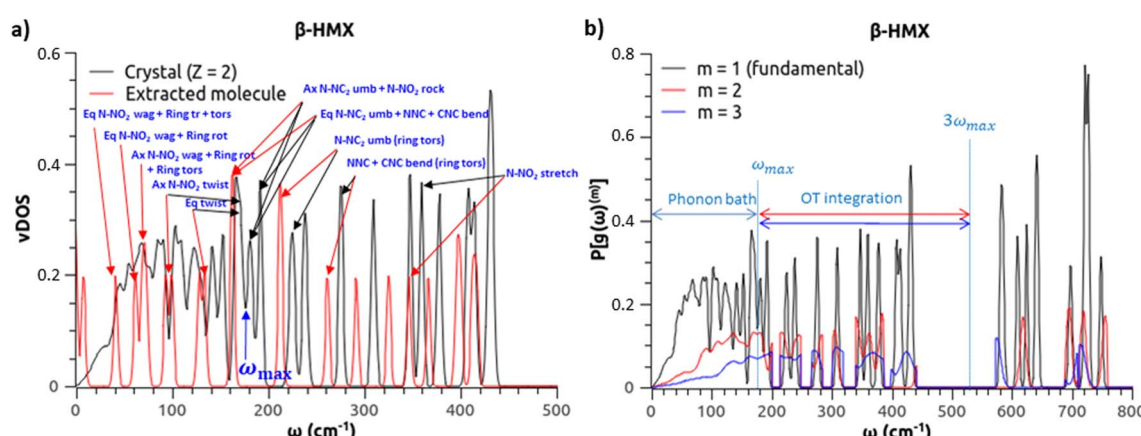


Fig. 1 (a) vDOS of the β -HMX molecular crystal (2 molecules per unit cell) and of the extracted HMX molecule in gas phase. The axial (Ax) and equatorial (Eq) N-NC_2 umbrella modes of the isolated molecule degenerate in the crystal. On the contrary, the N-NO_2 twisting modes of the Ax and Eq nitro groups of the isolated molecule synchronize in the crystal (large peak between 159 and 176 cm^{-1}). Since there is no N-NO_2 twisting modes beyond 176 cm^{-1} , this value defines ω_{\max} . (b) Fundamental vDOS (black) and projected OTs for $m = 2$ and $m = 3$ (red and blue, respectively), and integration domain. Using eqn (1) and (2), the resulting SSI for β -HMX is 0.9020.



study with ω_{\max} assigned to some other modes is out of the scope of the present work.

Then, the phonon density of the m^{th} OT ($m = 2, 3, \dots$) has the form $g(\omega)^{(m)} = g(\omega)/m$, using stretched frequencies $\omega^{(m)} = m\omega$. The projection onto the fundamental vDOS yields the quantity $P[g(\omega)^{(m)}]$, which is integrated from ω_{\max} to $3\omega_{\max}$ and normalized by the number of states in the phonon bath ($N_{\text{phonon bath}}$). The resulting shock sensitivity index is hereafter denoted SSI (eqn (1)). An example is provided for β -HMX in Fig. 1(b). Eqn (1a) yields the number of states in the phonon bath, which is $N_{\text{phonon bath}} = 30$ states. The contributions to eqn (1b) are 0.4707 and 0.4313 for $m = 2$ and $m = 3$, respectively, yielding a total SSI of 0.9020 for β -HMX. Michalchuk *et al.*³² have shown that only considering OTs $m = 2$ and $m = 3$ and adding up their contributions is enough to relate fairly well to h_{50} as obtained from mass drop experiments. This corroborates Bondarchuk's recent choice⁴⁴ to use a damping factor which attenuates the contribution of higher overtones.

Experimental h_{50} values can be found in Table 1. All of them but α -FOX-7's are from experiments using a hammer with grit, which enables significantly less uncertainty than a bare hammer, as statistically observed by Marrs *et al.*⁴⁵. They also observed that ignition and burning with a bare hammer are more inhomogeneous. For these reasons, tests with a gritted hammer are a better proxy to relate experimental h_{50} values to the present theoretical model, *i.e.*, to relate actual initiation to an initial vibrational up-pumping scheme.

$$N_{\text{phonon bath}} = \int_0^{\omega_{\max}} g(\omega) d\omega \quad (1a)$$

$$\text{SSI} = \sum_{m=2,3} \int_{\omega_{\max}}^{3\omega_{\max}} P[g(\omega)^{(m)}] d\omega / N_{\text{phonon bath}} \quad (1b)$$

III. Results and discussions

A. Prediction of h_{50} for various molecular crystals

1. Relationship between the shock sensitivity index SSI and experimental h_{50} . Data resulting from the OT-based model are displayed against experimental h_{50} in Fig. 2(a), for the molecular crystals of Table 1, which spread across a large range of sensitivity. It shows that the contribution of the two first overtones is relevant for the total to fit close to the black dashed line, from which high-sensitive β -CL20 and PETN deviate, as well as insensitive α -FOX-7. For the two formers, we suppose that the impact test under these standard conditions overdrives their sensitivity. A new standard with a smaller sample mass could resolve this, as discussed in Section III.B, or the use a different protocol.⁵⁵ For the latter, it is not the first time it behaves as an outlier.⁵⁶ This is also the only one in Table 1 tested with a BAM apparatus, which involves a bare 2 kg hammer and a calibrated sample volume instead of mass. Additionally, FOX-7 is the smallest molecule of this set and it may obey to a different mechanism. It should be noted that if we limit α -FOX-7's phonon bath to the highest libration mode instead of including the highest C-NO₂ twisting modes, its ω_{\max} is lowered to 139 cm⁻¹, yielding an SSI of 0.7500, which restores the agreement with the global trend.

Discarding α -FOX-7, a fit with an exponential decay function succeeds in modeling the trend, and all the SSI data align well

Table 1 Theoretical \widehat{h}_{50} from eqn (2) and experimental h_{50} values of various molecular crystals

	\widehat{h}_{50} (cm)	h_{50} (cm)	Ref.	Hammer (kg)	Tool type ^a	Sample mass (mg)	Remarks
PETN	12.6	13.8	45	2.5	12	40	Average of many h_{50} from only LANL
β -CL20	11.8	14	46	2.5	12	35	
ε -CL20	21.9	17.7	46	2.5	12	35	Average of the reported values
BTF	29.6	21	47	2.5	12	—	Non-recryst, highest dens. sample
β -HMX	32.6	32	48	2.5	12	35	33 cm with 12B and 2.0 kg (ref. 49)
δ -HMX	19.7	18	50	2.5	12	40	Av. of pure δ and δ with β traces
α -NTO	141	291	51	—	12	—	293 cm with tool 12B
HNB	21.5	15.6	52	2.5	12	30–40	
Tetryl	83.0	38.5	52	2.5	12	30–40	
HNAB	26.1	32	36	2.5	12	40	
HNS	48.8	39	36	2.5	12	40	HNS-II (denser than HNS-I)
DIPAM	44.0	85.1	36	2.5	12	40	
TNB	81.8	100	36	2.5	12	40	
MATB	119	177	36	2.5	12	40	
DATB	434	320	36	2.5	12	40	
TATB	585	490	36	2.5	12	40	>320 cm. Value extrapolated from TNB, MATB and DATB
α -FOX-7		126	53	2.0	BAM	—	Recrystallized sample, bare hammer
			54				
α -CL20		20.7	55	5.0	12	50	
β -CL20		24.2	55	5.0	12	50	
γ -CL20		24.9	55	5.0	12	50	
ε -CL20		26.8	55	5.0	12	50	

^a Tool types 12 and 12B: hammer with and without sandpaper, respectively.



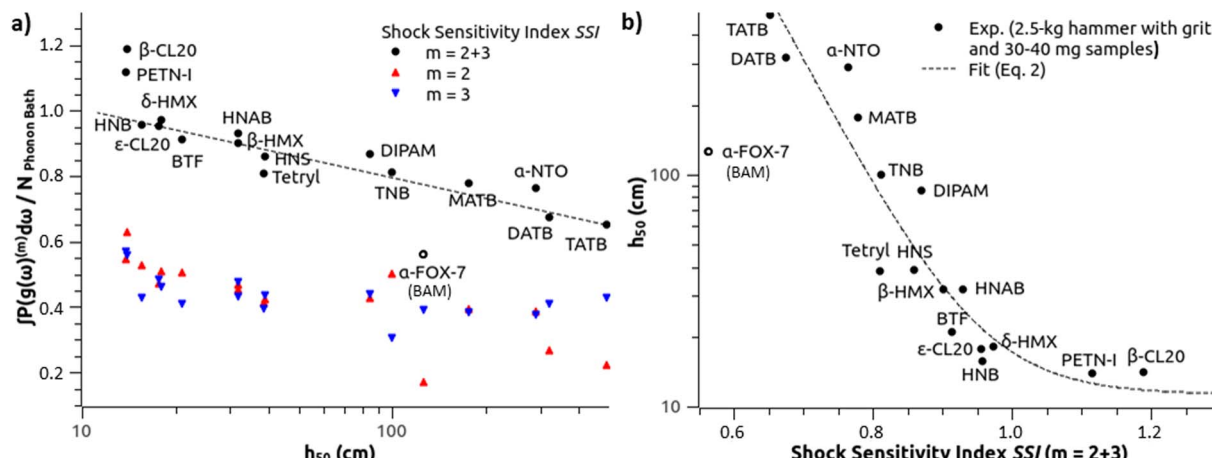


Fig. 2 SSI from the OT-based method to predict impact sensitivity of (a) various molecular crystals (Table 1), (b) the same with swapped axis. Red, blue and black color code is for OTs $m = 2$, $m = 3$, and total contributions, respectively. The dashed line in (a) is a visual guide from which two high-sensitive explosives and α -FOX7 deviate. The dashed curve in (b) displays the estimate \hat{h}_{50} determined as in eqn (2) (excluding α -FOX7 – see Section III.A.1).

along the black dashed curve representing the estimate \hat{h}_{50} in Fig. 2(b) (eqn (2) with $y_0 = 11.27$ cm, $a_1 = 509.58$ cm, $b_1 = 0.0761141$, and $SSI_0 = 0.660437$). The theoretical \hat{h}_{50} for the molecular crystals of interest are reported in Table 1. Using a log scale, the correlation coefficient R^2 of the OT-based model is 90%.

$$\hat{h}_{50}(\text{cm}) = y_0 + a_1 \exp\left(-\frac{SSI - SSI_0}{b_1}\right) \quad (2)$$

2. Sensitivity ranking of polymorphs

(a) **HMX.** β -HMX is one of the most investigated molecular explosives and experimental values of h_{50} are available.^{48,49,52} A few values for δ -HMX have also been reported, for instance for a pure sample and for some samples with traces of β -HMX.⁵⁰ Averaging them, the resulting value of 18 cm compares well with the 19.7 cm from the OT-based model (eqn (2)). The sensitivity of δ -HMX appears thus similar to that of ϵ -CL20.

However, there is still no consensus whether the observed higher sensitivity of δ -HMX comes from the molecular conformation or from the cracks generated by the β -to- δ phase transformation. Decorrelating the effects of porosity, because the particles were confined in a polymer matrix, δ -HMX was experimentally found to be intrinsically more sensitive than β -HMX and comparable to PETN.²⁷ Under different conditions, using a BAM apparatus and 10 mg samples, a recent experimental study⁵⁷ contradictorily claims that the molecular conformation of HMX has little to no relevance. Their figures show very similar pore/crack distributions in “small” particles of both pristine β -HMX and δ -HMX, which represents an opportunity to compare their intrinsic sensitivity. But the authors strangely deem that δ more sensitive than β by 2.6 cm is statistically not meaningful. Yet, this value corroborates the previous experimental finding as well as our theoretical observation. To emphasize the importance of molecular conformation and packing, quantum molecular dynamics simulations show that the onset of shock chemistry occurs at 2200 K for β -

HMX and 2000 K for δ -HMX, and involving different decomposition mechanisms.⁵⁸

Pristine β -HMX was transformed to δ -HMX and then reverted. Concatenating the recent results⁵⁷ with previous studies,^{27,59} a trend rather appears, as shown in Table 2. The smaller the initial particle size of pristine β , the less sensitive reverted- β is. But the “small” and “large” particle regimes involve different phenomena. In the “small” particle regime, the reverted- β particles look smoother than in pristine β , which is likely at the origin of their reduced sensitivity. In the “large” particle regime, reverted- β has crack porosity larger than in pristine β , explaining their higher sensitivity.

Some other factors affecting the sensitivity are addressed in Section III.B. Therefore, eqn (2) should be understood as a transfer function between the SSI and experimental h_{50} , for samples with the “standard” granulometry and porosity of the compounds displayed in Table 1.

(b) **CL-20.** At the impact and shock timescales, all CL-20 phase transformations but one ($\gamma \leftrightarrow \zeta$) are too slow to occur, and there is no liquid phase.⁶⁰ This makes the discussion for CL-20 more straightforward than for HMX. An experiment using a standard protocol and a hammer of 2.5 kg (ref. 46) ranks β -CL20 as more sensitive than ϵ -CL20, and our theoretical determination in Fig. 2(b) agrees well with the reported h_{50} values. Using another standard protocol and a hammer of 5.0 kg,⁵⁵ these values for α -, β -, γ - and ϵ -CL20 are measured at 20.7 cm, 24.2 cm, 24.9 cm and 26.8 cm, respectively. Fig. 3 compares

Table 2 Sensitivity of reverted- β vs. pristine β . The smaller the initial particle size of pristine β , the less sensitive reverted- β is

Initial particle size (μm)	$h_{50}[\text{rev-}\beta] - h_{50}[\beta]$ (cm)	Ref.
7–40	+7.3	57
100	+1.2	59
30–120	Same observed ignition behavior	27
600	-19	57



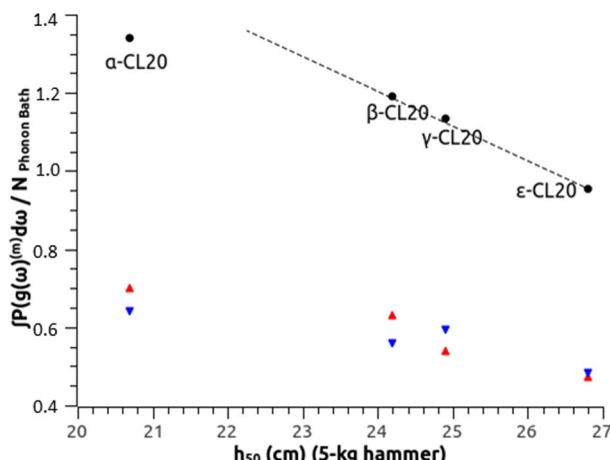


Fig. 3 SSI from the OT-based method to predict impact sensitivity of CL20 polymorphs (Table 1). The dashed line is a visual guide, which shows the alignment between β -, γ - and ϵ -CL20.

these determinations with the SSI, showing that the OT-based model agrees with the experimental ranking. Indeed, β -, γ - and ϵ -CL20 are visually well aligned along the black dashed line, which supports once more the effect of molecular conformation and packing on sensitivity. α -CL20 appears a bit under, but it always naturally contains small molecules like H_2O ,⁶¹ CO or CO_2 ,⁶⁰ which could affect its effective sensitivity. Using eqn (2), our estimates for 35 mg samples tested with a 2.5 kg grit hammer would be 11.3 cm, 11.8 cm, 12.3 cm and 21.9 cm, respectively. It may be difficult to experimentally discriminate α -, β -, and γ -CL20 with this standard protocol, due to overdriven sensitivity, as mentioned in Section III.A.1.

3. Sensitivity ranking cocrystals and their counterparts.

Impact tests using the standard BAM method were performed for ϵ -CL20, MTNP and the 1:1 cocrystal of CL20:MTNP,⁶²

yielding h_{50} values of 20.4 cm, 125 cm and 30.6 cm, respectively. The sensitivity of the cocrystal is expected to be similar to that of β -HMX. From eqn (2), our estimates for 35 mg samples tested with a 2.5 kg grit hammer are 21.9 cm, 444 cm and 11.8 cm, respectively. If MTNP is successfully predicted as insensitive, the cocrystal is wrongly predicted as highly sensitive. We have checked for configurational issues in the cocrystal structure. In the cocrystal, the CL20 molecules are in the γ conformation as expected. When only considering the number of doorway modes, a recent study found the right ranking.⁶³ But Michalchuk *et al.*³² have demonstrated that this simple approach is more qualitative than quantitative, and lacks transferability over a more complete set of molecular crystals.

Fig. 4(a) compares the theoretical SSI to experimental h_{50} of ϵ -CL20, β -HMX and the 2:1 cocrystal, as measured by Bolton *et al.*,¹⁴ using very small amounts of explosive (0.5 mg) and a hammer of 2.5 kg. Here again, the cocrystal is wrongly predicted as very sensitive. Note that instead of a random spatial distribution of the CL20 molecules in β or γ conformations in the cocrystal at ambient conditions, our optimized cell only contains CL20 molecules in γ conformation. Starting the optimization from CL20 molecules all in β conformation ends up with all in γ conformation. Even though it is not certain how the presence of CL20 molecules in β conformation would affect the sensitivity, the model fails to rank the sensitivity of the cocrystals with respects to their counterparts.

The model needs a correction for cocrystals. Long range and local couplings between modes are presumably more complex due to a higher number of possible combinations involving van der Waals interactions, hydrogen bonds of various intensities, or some weaker intermolecular $\text{O}\cdots\text{O}$, $\text{O}\cdots\text{N}$ and $\text{N}\cdots\text{N}$ interactions.⁶⁴⁻⁷⁰ Note that despite individually weaker than hydrogen bonds, the cumulative population of $\text{O}\cdots\text{O}$ and $\text{O}\cdots\text{N}$ interactions makes them the prevailing intermolecular contribution in ϵ -CL20.⁶⁹ Some of these easily breakable $\text{O}\cdots\text{O}$ bonds

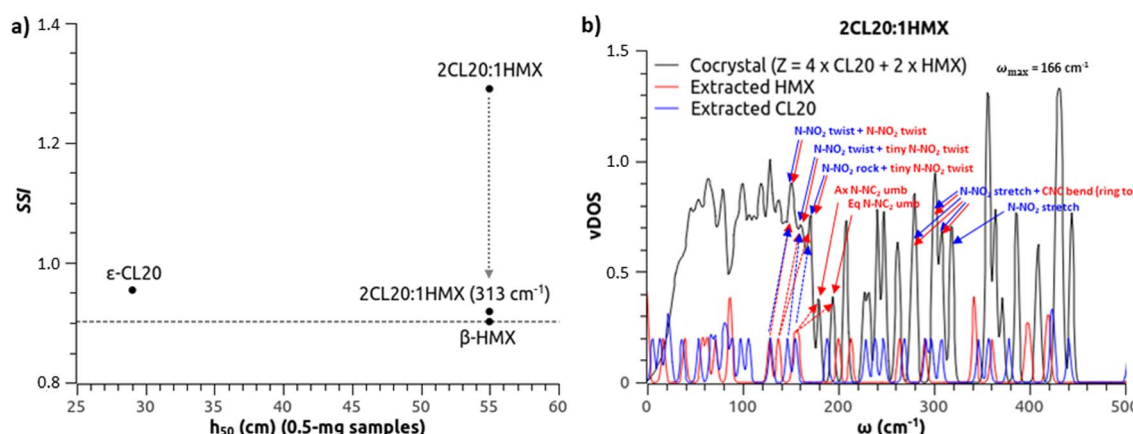


Fig. 4 (a) SSI from the OT-based model to predict impact sensitivity of 2CL20:1HMX cocrystal and its counterparts. Using the same phonon bath criterion as mono-molecular crystals, the model fails to rank the cocrystal between its counterparts. Additionally amalgamating to the phonon bath the modes before a non-coupled N-NO_2 stretching mode restores the agreement ($\omega_{\text{max}} = 313 \text{ cm}^{-1}$). (b) vDOS of the cocrystal (black) and the extracted molecules of HMX (red) and CL20 (blue) in gas phase, and some phonon assignments. Note how the lowest N-NO_2 twist modes shown for extracted HMX and CL20 (129 cm^{-1}) stay synchronous in the cocrystal (151 cm^{-1}), despite shifted. Note how the next N-NO_2 twist mode in extracted HMX synchronizes to the N-NO_2 twist and rock modes of extracted CL20 in the cocrystal. Note also how the axial and equatorial N-NC_2 umbrella modes in extracted HMX degenerate in the cocrystal (like in β -HMX).



could be advantageously replaced in the vicinity of a co-molecule like HMX, involving a coupling of the pertaining moieties. While co-crystals are expected to be of lower sensitivity if the most sensitive component is mixed with a lower sensitivity molecular partner, the up-pumping mechanisms can be significantly different. The co-molecules are probably more coupled than expected (see a few mode assignments in Fig. 4(b)) and a broader phonon bath could make sense. For instance, considering the highest non-coupled N-NO₂ stretching mode and selecting the coupled N-NO₂ stretching modes before (CL20 vibrations synchronized to HMX ring torsion (Fig. 4(b)) or to MTNP ring twisting, respectively) and amalgamating them to the phonon bath, we find $\omega_{\max} = 313 \text{ cm}^{-1}$ for 2CL20 : 1HMX and $\omega_{\max} = 311 \text{ cm}^{-1}$ for 1CL20 : 1MTNP. This yields SSI = 0.9189 for 2CL20 : 1HMX and SSI = 0.9739 for 1CL20 : 1MTNP. This ranks 2CL20 : 1HMX slightly more sensitive than β -HMX, as expected, and 1CL20 : 1MTNP similar to ϵ -CL20, which is better but still a bit high.

Sensitivity prediction for cocrystals seems more subtle and complex than for mono-molecular crystals. Up-pumping schemes for cocrystals should also probably include energy transfer rates. Looking at N-NO₂ stretching modes (Fig. 4(b)), which could trigger bond rupture, the corresponding lowest frequencies are the peaks at 280, 301 and 308 cm⁻¹. Dividing these values by 2, the resulting frequencies fall in the part of the phonon bath involving N-NO₂ twisting in both CL20 and HMX molecules. Therefore, a vibrational cooling of the N-NO₂ stretching modes by the phonon bath is very likely, depending on the specific mode-to-mode scattering rates, in turn reducing the sensitivity. This vibrational cooling was recently demonstrated to theoretically happen in α -RDX.³⁸

B. Experimental uncertainty and factors affecting the relationship with the SSI

The relationship between the SSI and the estimate \widehat{h}_{50} (eqn (2)) relies on experimental data. For consistency, we used data measured from similar tool type, hammer weight and sample mass (Table 1). But even for this consistent set, we must keep in mind that experimental errors can be as large as 20%. For instance, 435 tests on PETN performed at LANL over the past decades have been recently analyzed by Marrs *et al.*⁴⁵ to investigate various sources of sensitivity variations. They were able to discard factors like when was and who performed the measurements, because of no substantial effect. All tests were performed using a 2.5 kg hammer. The resulting mean h_{50} value and 95% confidence interval are $13.8^{+3.2}_{-2.5}$ cm with grit (hammer with sandpaper, like tool type 12 in Table 1) and $23.0^{+5.3}_{-4.7}$ cm without it (bare hammer, like tool type 12B). This represents around 20% of maximum experimental uncertainty (7% on a log scale). Moreover, the difference with/without grit is as large as 9.2 cm, which is very significant. Discrepancies between laboratories also exist, with -4.4 cm on average (more sensitive) in the cumulated labs other than LANL. The various "Go/No go" methods have less effects, although the discrepancies are within 2.9 cm. Nevertheless, they demonstrate that this latter factor has significant effects on variability of TNT

sensitivity. Uncertainty is expected to increase with less sensitive materials, supporting the use of a log scale. They conclude that, while ranking explosives in the range 8–12 cm is not reliable, relative observations performed in the same lab under the same conditions remain true.

Considering the granulometry of the samples, sensitivity predictions can become less intuitive. In the same paper,⁴⁵ the h_{50} values of PETN comprised of small ($\approx 26.08 \mu\text{m}$) or large ($\approx 331.7 \mu\text{m}$) particles are measured. With a bare hammer, small particles are found less sensitive than large ones (by roughly 3 cm) as expected,^{71,72} but it is counterintuitively the opposite with grit (by roughly 3 cm). A similar apparent inconsistency between impact sensitivity and grain size of HNS pressed to a single density was shown to depend on the shock amplitude and duration,⁷³ and the author mentioned a similar behavior for RDX and TATB. The non-monotonic sensitivity with median micrograin size of 10%-porous RDX was also shown.⁷⁴ This non-monotonic behavior was theoretically confirmed for HMX, TATB and TNT.⁷⁵ At a given porosity, the voids must be sufficiently small, depending on the considered explosive, for the sensitivity to decrease and ignition to eventually fail. From reactive molecular dynamics, decreasing the size of nanoparticles in nanostructured RDX have been found to inhibit the shock-induced chemistry.⁵

The crystal morphology also affects its sensitivity. The behavior of "single" (S) or "multiple" (M) crystals, as named by Cole in his study⁴⁹ (the latter can be understood as powders), is different. His h_{50} values are reported in Table 3. Without grit, it seems that the cyclic nitramines (HMX and RDX) are barely affected by the crystal morphology, whereas the nitramine PETN behaves as expected ((M) more sensitive than (S)), and in an opposite way for nitroaromatic TNT. With grit, the sensitivity variations with the sample mass make the trends less clear: the sensitivity decreases as the sample mass increases, except for the single crystal of TNT and the powder of HMX. The exception is confirmed for the latter with an h_{50} value as low as 55 cm for a tiny 0.5 mg sample of β -HMX powder.¹⁴ Extrapolating, there must be for HMX a sample mass threshold for which (S) becomes more sensitive than (M), and the opposite for TNT.

To complete the picture, shock-induced polymorphic transitions are likely to occur more than anticipated. This is the case for γ -FOX-7, which reverts to α -FOX-7 during drop-weight experiments.⁴³

Table 3 Summary of experimental h_{50} values of 20 mg "single" (S) and "multiple" (M) crystals as measured using a 2 kg hammer without/with grit⁴⁹

h_{50} (cm)	Without grit		Sensitivity (S) vs. (M)	With grit		Sensitivity (S) vs. (M)
	(S)	(M)		(S)	(M)	
PETN	33	15	—	10 ^a	10 ^a	=
HMX	33	33	=	16 ^a	20 ^b	+
RDX	43	46	≈	18 ^a	30 ^a	+
TNT	53	102	+	56 ^b	25 ^a	—

^a Sensitivity increases (h_{50} decreases) when sample mass decreases.

^b Sensitivity decreases (h_{50} increases) when sample mass decreases.



In the light of all these experimental variabilities, we can understand why the sensitivity models presented in the introduction, fitted on raw and patchy h_{50} data, barely outperform the 80% accuracy. Gathering h_{50} data of dozens or hundreds of explosive compounds and, whether using machine learning or not, connecting these data to simple molecular, thermodynamic or kinetic descriptors can help understand the molecular origins of trends. But to take a step further in impact sensitivity engineering, it is recommended to thoroughly take into account the factors addressed above, which are often overlooked. Improved experiments may systematically report the sample granulometry/porosity and morphology, or define standards for these factors as well, which could help machine-learned models sort out the role of these factors.

Providing more than trends, the OT-based model can discriminate the sensitivity of the polymorphs of the same molecule. This feature is made possible due to the use of the phonon spectrum, which intrinsically contains more information than molecular descriptors. The vDOS is like a fingerprint of both the molecule and the crystal. Additionally, the smart post-processing proposed by Michalchuk *et al.*^{32,33} in their method gives an insight of the underlying physics. The present agreement of \widehat{h}_{50} validates the indirect up-pumping mechanism occurring under these conditions, with the doorway modes also interacting with the phonon bath during the shock initiation stage (Section II.C). A more detailed approach of the mechanism and straight overtone relationships have been used by Bondarchuk⁴⁴ in his recently revisited impact theory. Of course, once the hotspots are initiated or melting begins, further activation may not follow the doorway mode ordering.⁷⁶

The estimate \widehat{h}_{50} is fitted on a subset of experimental data whose maximum uncertainty is evaluated at 7% on a log scale.⁴⁵ The accuracy of the OT-based model is very good ($R^2 \approx 90\%$ on a log scale), and we believe that this great achievement is due to the use of high-quality DFT-D to perform the phonon calculations. Nonetheless, we want to emphasize that the predicted \widehat{h}_{50} (eqn (2)) is valid only for experiments performed under conditions similar to that of Fig. 1(b).

The theoretical SSI is determined for bulk crystals. Its relationship with the derived \widehat{h}_{50} may change, depending on the subset of h_{50} data it is adjusted on. For instance with a subset of h_{50} data pertaining to the bare hammer only, it is likely that the phonon bath extent ω_{\max} should be adjusted with a less stringent criteria for amalgamated intramolecular modes. This will be the subject of further investigations in future to drive studies on shock-resilient energetic crystals.

IV. Conclusion

In order to devise an accurate predictive model for impact sensitivity of molecular crystals, we used the promising overtone (OT) based approach by Michalchuk *et al.*³² in conjunction with high-quality phonon calculations. For the determination of the phonon bath extent, we systematically amalgamated the highest X-NO_2 twisting modes. The resulting intrinsic shock sensitivity index (SSI) was compared to the most common marker of impact sensitivity, h_{50} , as determined from drop

weight impact tests. When the data subset is selected from experiments performed under very similar conditions (2.5 kg grit hammer and 30–40 mg samples), the model can predict h_{50} values with very good accuracy. Eqn (2) acts like a transfer function between the theoretical SSI and experimental h_{50} . The parameters of eqn (2) do not explicitly account for overlooked aspects like sample granulometry or porosity. Eqn (2) thus applies to the “standard” granulometry and porosity of the compounds reported in Table 1.

This model has the ability to discriminate polymorphs, ranking δ -HMX as more sensitive than β -HMX, and correctly ranking the four polymorphs of CL20 as well. This very good agreement for mono-molecular crystals validates the initial indirect up-pumping mechanism occurring under these conditions, where the doorway modes also interact with the phonon bath.

However, the case of the cocrystals is more subtle and the phonon bath criterion used for mono-molecular crystals does not transfer well. Owing to the vibrational coupling of the molecules, it seems a broader phonon bath can be more appropriate. This also renders the choice of ω_{\max} less straightforward and intuitive, although our physics-based approach showed that frequencies of about 310 cm^{-1} could be considered.

Recalling experimental uncertainty and how various experimental factors affect h_{50} values for a given molecular crystal, we suggest the sample granularity, porosity, mass and morphology be systematically considered and reported with measurements. This will in turn allow sharpening the predictive capabilities of sensitivity models.

The OT-based model has a calculation cost. Accelerating phonon calculations without compromising accuracy in the low-frequency range pertaining to lattice and intermolecular vibrations becomes essential. This may be achieved using machine learning techniques, as successfully done for inorganic materials.^{77–79} These techniques have shown promising progress for molecular crystals.^{80,81} Meanwhile, the OT-based model could predict the h_{50} value of computationally-designed mono-molecular crystals with engineered sensitivity, as if they were tested with a 2.5 kg hammer with grit and using 30–40 mg samples.

Data availability

The data that support the findings of this study are available within the article and its ESI.†

Conflicts of interest

The authors have no conflicts to disclose.

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