1 Introduction

The co-crystallization strategy is being pursued in pharmaceuticals and optical materials. Later, the strategy has been introduced into the field of energetic materials. Co-crystallization can change the way of molecular assembly and arrangement, resolving the contradiction between energy density and safety to a certain extent, which has attracted wide attention from scholars. In 2011, Bolton and Matzger first synthesized a co-crystal material with a molecular number of 1:1 based on CL-20 with high energy density, high sensitivity and TNT with low energy density, low sensitivity and low cost. Its various properties are between the two elemental components, significantly reducing the sensitivity of CL-20. Since then, scholars in the field of energetic materials have successively synthesized a variety of CL-20-based co-crystals materials. Co-crystallization strategy has expanded new directions for the development of high-energy insensitive energetic materials.

To further explore the fundamental ways of co-crystallization strategy to improve material properties, scholars have carried out a lot of research work using theoretical analysis and simulations. Zhang et al. analyzed the energy characteristics and safety of 41 kinds of energetic co-crystals using theoretical calculations, obtaining common factors affecting the performance of co-crystals, which provided support for the design of new materials. They also analyzed the influence of crystal packing on molecular stability through quantitative calculations. The results showed that the co-crystallization can increase the interaction between molecules or improve the crystal packing mode, reducing shock sensitivity. Based on density functional theory calculations, Liu et al. analyzed the behavior of the two components in CL-20/HMX co-crystals under high pressure. Xue et al. used the molecular dynamics method based on the ReaxFF force field to compare the thermal decomposition processes between the CL-20/HMX co-crystals and single crystals of two components. The study believed that co-crystallization was achieved by changing the internal stability and intermolecular interaction to improve overall stability. Liu conducted a simulation study on the thermal decomposition process of the CL-20/TNT co-crystals, and later he also explored the process of reaction flow propagation in the CL-20/TNT co-crystals. Guo compared the thermal decomposition processes of the CL-20/TNT co-crystals, single crystals of the two components and a simple physical mixing system. The results showed that co-crystallization can well reconcile the two components. At the same time, the energy release of the co-crystals in the early stage of the reaction is slower than that of the physical mixing system. By analyzing the thermal
decomposition processes of two typical co-crystals of CL-20/TNT and CL-20/HMX, Ren summarized the three stages of thermal decomposition of CL-20-based co-crystals. In the actual production, storage, transportation and use of energetic materials, in addition to the thermal action from the outside, they are also likely to be shocked. A lot of theoretical research has been carried out on this aspect. He et al. used the SCC-DFTB (self-consistent charge density functional tight binding) method to study the initial decomposition mechanism of HMX and TATB under shock. Liu’s research on the reaction process of TNT under shock shows that the formation of TNT dimer has a great influence on shock sensitivity. Huang studied the anisotropy of shock sensitivity of insensitive explosive TATB, finding that the unique intermolecular hydrogen bond and different compressibility are the main reasons for the anisotropy of shock sensitivity. Xue used molecular dynamics to study the effect of dislocations on the shock sensitivity of RDX. Later, a similar method was used to study the initial decomposition mechanism of CL-20 under the action of shock waves with velocities of 8–11 km s\(^{-1}\). The results show that the shock wave velocity has a great influence on the reaction path, products and cluster formation. Liu studied the initial decomposition mechanism of CL-20/HMX co-crystals under a steady shock wave. Zhang used quantitative calculations to study the initial decomposition process of CL-20/TNT co-crystals under shock. The above researches are mainly based on the balance impact process simulation method of multi-scale shock technology. Other studies have simulated the response process of energetic materials such as TATB, PETN, RDX, HMX and ICM-102 by the non-equilibrium impact method.

A large number of theoretical studies on energetic co-crystals mainly focus on the analysis of its thermal decomposition process, the comparative understanding of the thermal decomposition process of co-crystals and single crystals of their components is relatively sufficient. People also have a deep understanding of the shock response of energetic single crystals. Nevertheless, there are relatively few studies on the response process of energetic co-crystals under shock, especially the anisotropy of shock response caused by the layered structures of co-crystals that have not been reported. In this work, the anisotropy of the response of the typical energetic co-crystals CL-20/TNT under shock is studied theoretically, to provide a reference for the understanding of the shock sensitivity of energetic co-crystals and the design of new energetic co-crystals materials.

2 Methods and computational details

Co-crystals cell data on CL-20/TNT used in this work were derived from the X-ray crystal structure. The initial single-crystal cell contains 8 CL-20 molecules and 8 TNT molecules. Based on this, the single-crystal cell was expanded to a \(4 \times 2 \times 1\) supercell, which contains 64 CL-20 molecules and 64 TNT molecules, totaling 3648 atoms.

Firstly, the conjugate gradient algorithm was used to relax the co-crystal structures. The convergence tolerance of force was \(10^{-7}\) kcal mol\(^{-1}\) Å\(^{-1}\). Subsequently, we carried out a canonical ensemble (NVT) MD simulation each for 10 ps at 298 K using the Berendsen thermostat to relax the supercell. To obtain the structure at atmospheric pressure, the NPT ensemble was used for 15 ps relaxation at 298 K and 0 GPa, the Nosé–Hoover method was selected for temperature and pressure controls. The co-crystals structure of CL-20/TNT at room temperature and pressure was obtained, the density was 1.89 g cm\(^{-3}\). Then, steady shock waves with velocities of 7, 8 and 9 km s\(^{-1}\) were loaded along X, Y and Z using multi-scale shock technology.
There were 9 simulations with 0.1 fs timestep, periodic boundary conditions and total durations of 50 ps. All simulations use the Lammps package, the potential function was ReaxFF/lg.\textsuperscript{12} The action directions of shock waves are shown in Fig. 1.

3 Results and discussion

3.1 Evolution of temperature, stress and volume

The temperature evolution of the co-crystal system under shock-wave loading along different directions with different velocities is shown in Fig. 2. The temperature rose under various conditions. Under the shock wave with a velocity of 7 km s\textsuperscript{-1}, the temperature rose sharply in the initial stage and then remained stable. Under the shock wave with a velocity of 8 km s\textsuperscript{-1}, the temperature rose sharply in the initial stage and then rose at a slower rate. Under the shock wave with a velocity of 9 km s\textsuperscript{-1}, the rise of the temperature underwent 3 stages: rose sharply, rose rapidly and rose slowly. In the initial stage, the shock wave led to a physical change in the system, resulting in a sharp temperature rise. At 7 km s\textsuperscript{-1}, the rising temperature could not lead to a certain scale chemical reaction, making the temperature remain stable. At 8 km s\textsuperscript{-1}, the rising temperature could lead to a certain scale chemical reaction, making the temperature rise at a slower rate. At 9 km s\textsuperscript{-1}, the rising temperature could lead to a large-scale chemical reaction, making the temperature rise rapidly. After that, the chemical reaction rate slowed down and the system temperature rose at a slow rate. Comparing changes of the system temperature caused by the shock wave with the same velocity in different directions, it can be found that the shock wave loaded along the $Y$ direction makes the temperature of the system higher than that along the $X$ and $Z$ directions. It shows that the shock temperature response of CL-20/TNT co-crystals has a certain anisotropy, which is caused by its special layered structure.

The stress evolution of the co-crystal system under shock-wave loading along different directions with different velocities is shown in Fig. 3. Comparing changes of the system stress caused by shock waves along the same direction with different velocities, it can be seen that the shock wave with higher velocity led to higher stress. Comparing the changes in the system stress caused by shock wave loading along different directions with the same velocity, it was found that the shock wave loaded along the $Y$ direction made the system stress significantly higher than that along the $X$ direction and $Z$ direction. This shows that the shock stress response of CL-20/TNT has a certain anisotropy, which is also related to its special layered structure.

The volume evolution of the co-crystal system under shock-wave loading along different directions with different velocities is shown in Fig. 4. The value given in the figure is the ratio of the current volume to the volume before the shock wave, which reflects the degree of compression. It can be observed from the figure that the shock wave will cause the system to be compressed under all conditions. Along the same direction, the shock wave with higher velocity led to a higher degree of compression. With the same velocity, the volume change curves caused by shock waves along different directions are similar, but the compression degree along the $Y$ direction is significantly higher than that along $X$ and $Z$ directions, that is, the $Y$ direction is easier to be compressed.

Zhang's research found that the higher degree of compression would lead to the higher temperature and stress of the CL-
According to the above analysis of temperature, stress and volume evolution information, it is found that there are obvious differences between the shock wave loaded along the Y direction and X, Z directions. The Y direction is easier to be compressed, resulting in higher temperature and stress in the system, which will lead to obvious differences in the process of the reaction.

3.2 Reactants and products

Fig. 5 shows the amount evolution curves of reactants and products under various conditions. Firstly, we analyze the decomposition of reactants CL-20 and TNT. When the shock wave with higher velocity was loaded, the reactants decomposed more rapidly and completely. Under the shock wave with a velocity of 7 km s\(^{-1}\), the reaction degree of the whole system was very low, CL-20 and TNT did not decompose completely within 50 ps. Under the shock wave with a velocity of 8 km s\(^{-1}\), the decomposition rate of reactants was accelerated, reactants decomposed completely within 50 ps. Under the shock wave with a velocity of 9 km s\(^{-1}\), the reactants almost decomposed completely within 50 ps.
completely in 10 ps. Comparing the effects of shock waves with the same velocity along with different directions, it can be observed that the decomposition rate of the system along the Y direction is the fastest, that along the Z direction is the second, and that along the X direction is the lowest.

Then, we analyze the main intermediates. Under the shock wave with a velocity of 7 km s\(^{-1}\), the reaction degree of the system was low, there was only a small amount of NO, NO\(_2\) and NO\(_3\) within 50 ps. Under the shock wave with a velocity of 8 km s\(^{-1}\), within 50 ps, there were certain amounts of NO, NO\(_2\) and NO\(_3\) in the system, indicating that the whole reaction continued at a certain rate. Under the shock wave with a velocity of 9 km s\(^{-1}\), the intermediate products NO, NO\(_2\) and NO\(_3\) were rapidly produced at the beginning and almost consumed within 20 ps, indicating that the reaction developed rapidly at the beginning, completed the large-scale reaction process in a short time, and then slowed down.

Finally, we analyze the final products. With the velocity of 7 km s\(^{-1}\), there was no final product along the X and Z directions. However, along the Y direction, due to the higher degree of compression, the temperature and stress were also higher, the reaction degree was improved, a small amount of N\(_2\) was produced. With a velocity of 8 km s\(^{-1}\), along the X direction there was a certain amount of N\(_2\) (63 at 50 ps), and a small amount of H\(_2\)O. Along the Z direction, there was more N\(_2\) (90 at 50 ps) and H\(_2\)O, there was even a small amount of CO\(_2\) in the system, indicating that the carbon rings in the system had begun to break and form small molecules containing carbon. Along the Y direction, there was the largest amount of N\(_2\) (115 at 50 ps) and H\(_2\)O, there was also a small amount of CO\(_2\) in the system. At the same time, there was a small amount of NH\(_3\). With a velocity of 9 km s\(^{-1}\), large-scale chemical reactions took place in the system in a short time. The final products were mainly N\(_2\), H\(_2\)O, CO, CO\(_2\) and NH\(_3\), which were significantly more than those at 8 km s\(^{-1}\). The differences between different directions were relatively small.

From the evolution of reactants and products of the system, it can be seen that the velocity of the shock wave can directly affect the decomposition of reactants, the types and number of products. The shock wave with higher velocity, made reactants decompose more rapidly and completely, leading to richer products. In addition, under the shock wave with the same velocity along different directions, the changes in reactants and products showed anisotropy. The shock wave along the Y direction can bring greater compression, greater stress and higher temperature, resulting in faster decomposition of reactants and richer stable products.

3.3 Cluster

3.3.1 Amount of cluster. The evolution of cluster amount is shown in Fig. 6. Under the shock wave with a velocity of 7 km s\(^{-1}\), the amount of cluster continued to rise. The amount along the Y direction was the most; that along the Z direction was the second. The shock caused the system to be compressed, the distance between atoms decreased, the probability of mutual collision increased, and clusters continued to be generated. Under the shock wave with a velocity of 8 km s\(^{-1}\), the amount of cluster rose rapidly at the beginning and gradually decreased after reaching the peak. The shock caused the system to be compressed rapidly, more clusters are produced in a shorter time. After that, chemical reactions occurred, part of the cluster gradually decomposed into smaller molecules. In the first increasing stage, shock waves along the Y direction induced greater compression of the system, there were more clusters generated. Nevertheless, in the next decreasing stage, shock waves along the Y direction induced more intense chemical reactions, there was more cluster decomposition. Under the shock wave with a velocity of 9 km s\(^{-1}\), the number of clusters...
rose sharply at the beginning, then decreased rapidly, finally fluctuating within a certain range. As previously analyzed, there were three stages of evolution: high compression stage, large-scale chemical reaction stage and stabilization stage. Due to the high intensity of shock waves, there were few differences in the action direction of shock waves.

### 3.3.2 Composition of cluster

Through analysis of the data, it can be noted that certain amounts of C_{12} and C_{13} appear in the cluster. CL-20 contains 6 carbon atoms in one molecule and TNT contain 7 carbon atoms in one molecule. That is, C_{12} is a cluster composed of two CL-20 molecules, and C_{13} is a cluster composed of one CL-20 molecule and one TNT molecule. Fig. 6 shows the changes in the amounts of C_{12} and C_{13} in the system. At 7 km s\(^{-1}\), along the X direction, C_{12} was generated at 1 ps, the total amount of C_{12} was the most with this velocity, C_{13} was generated at 4 ps. Along the Y direction, C_{13} was generated at 1 ps, the total amount of C_{13} was the largest with this velocity, C_{12} only existed in a small amount in the subsequent part of the time. Along the Z direction, C_{13} was generated at 1 ps, C_{12} also existed only in a small amount in the subsequent part of the time.

At 8 km s\(^{-1}\), along the X direction, there was a considerable amount of C_{12} in the system, and the peak amount of C_{13} was the lowest with this velocity. Along the Y and Z direction, a large amount of C_{13} was generated in the initial stage, their amounts gradually decreased after reaching the peak, there was a certain amount of C_{12}. At 9 km s\(^{-1}\), due to the violent chemical reaction in the system, the differences mainly existed before 10 ps. The sum of C_{12} and C_{13} along each direction had little difference. Along the X direction, the amount of C_{12} was more, and the corresponding amount of C_{13} was less. From the composition of the cluster under different shock waves, we can find that along the X direction, the probability of forming CL-20 dimers in the system is higher than that along other directions, and the probability of one CL-20 molecule binding to one TNT is relatively low. As shown in Fig. 1, due to the special layered structure of the co-crystals system, there will be great differences in the aggregation mode between molecules under shock waves along different directions, resulting in the anisotropy of the system responses.

### 3.3.3 Size distribution and mass ratio of cluster

Fig. 7 shows the distribution of cluster sizes in the system under different conditions. At 7 km s\(^{-1}\), along the X direction, the total amount of cluster was the least, most of the cluster was concentrated in C_{13}−C_{22}. Along the Y direction, the total amount of cluster was the most and most of the cluster was concentrated in C_{13}−C_{22}. Along the Z direction, the total amount of cluster was in the middle, most of the cluster was concentrated in C_{13}−C_{22}, but there was still a certain number of larger Z clusters. At 8 km s\(^{-1}\), the sizes of clusters along the X direction were small in the early stage, and the distribution along the Y direction was similar to that along the Z direction. At 9 km s\(^{-1}\), the shock wave intensity was high, and the differences caused by different directions were relatively small.

Due to the large differences in the sizes of clusters, the amount cannot comprehensively describe the overall situation of the cluster in the system. It is also necessary to analyze the mass ratios of the cluster. The mass ratio of the cluster is the ratio of the total mass of the cluster to the total mass of all atoms in the system. As shown in Fig. 8, at 7 km s\(^{-1}\), along the X direction the mass ratio of the cluster was the smallest.

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**Fig. 7** Cluster size distribution.

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**Fig. 8** Cluster size distribution.
In the response process was analyzed. The results show that: crystals under shock waves were simulated, and the anisotropy directions, respectively. The response processes of energetic co-crystals, when the shock waves are weak, the amounts, compositions, sizes and mass ratios of clusters show obvious anisotropy. When the shock waves are strong enough, the mass ratios in each direction were relatively close. At 9 km s$^{-1}$, as previously analyzed, the shock wave intensity was high enough, and the differences caused by different directions were relatively small (Fig. 9).

4 Conclusions

ReaxFF force field and multi-scale impact technology were used for generating shock waves with velocities of 7, 8, 9 km s$^{-1}$ that were loaded into CL-20/TNT co-crystals along with $X$, $Y$ and $Z$ directions, respectively. The response processes of energetic co-crystals under shock waves were simulated, and the anisotropy in the response process was analyzed. The results show that:

1) With the same velocity, compared with the shock wave along the $X$ direction and $Z$ direction, the shock wave along the $Y$ direction will make the compression degree of the system higher, resulting in higher temperature, higher stress and a higher degree of chemical reactions.

2) From the evolution of reactants and reaction products, along the $Y$ direction the reactants decompose more rapidly and the types of products are more abundant.

3) In terms of cluster evolution, due to the special structure of CL-20/TNT co-crystals, when the shock waves are weak, the amounts, compositions, sizes and mass ratios of clusters show obvious anisotropy. When the shock waves are strong enough, this anisotropy is not obvious.

Conflicts of interest

There are no conflicts to declare.

Notes and references

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