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OPEN Self-assembly of Carbon Vacancies in Sub-stoichiometric ZrC_{1-x}

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Sub-stoichiometric interstitial compounds, including binary transition metal carbides (MC1-x), maintain structural stability even if they accommodate abundant anion vacancies. This unique character endows them with variable-composition, diverse-configuration and controllable-performance through composition and structure design. Herein, the evolution of carbon vacancy (V_c) configuration in substoichiometric ZrC_{1-x} is investigated by combining the cluster expansion method and first-principles calculations. We report the interesting self-assembly of V_c s and the fingerprint V_c configuration (V_c triplet constructed by 3^{rd} nearest neighboring vacancies) in all the low energy structures of ZrC_{1-x} . When V_c concentration is higher than the critical value of 0.5 (x > 0.5), the 2nd nearest neighboring V_c configurations with strongly repulsive interaction inevitably appear, and meanwhile, the system energy (or formation enthalpy) of ZrC_{1-x} increases sharply which suggests the material may lose phase stability. The present results clarify why ZrC_{1-x} bears a huge amount of V_cs , tends towards V_c ordering, and retains stability up to a stoichiometry of x = 0.5.

Most covalent and ionic crystalline solids (daltonide) hold exact stoichiometry in order to keep translational symmetry and atomic coordination. In contrast, another group of compounds (berthollide, such as binary transition metal carbides and nitrides) maintain structural stability in a wide sub-stoichiometric range¹⁻³. These materials have rock-salt crystal structure (B1) with C/N atoms locating at the octahedral interstitial sites of the f.c.c. sublattice constructed by transition metal atoms. The interstitial atomic sites are easy to form high concentration of C/N vacancies. For example, the concentration of carbon vacancies accommodated in TiC_{1-x} and ZrC_{1-x} is as high as 50%^{4,5}. Both short-range ordering (SRO) and long-range ordering (LRO) of anion vacancy distribution are common in sub-stoichiometric materials⁶. Ordered phases can be fabricated by long-duration post annealing and rapid spark plasma (SPS) processing etc.⁷. These unique characters provide us the opportunity to succeed in defect engineering through modification of chemical composition and vacancy configuration. The related compounds were classified as 'non-stoichiometric interstitial compound'2, and have attracted extensive interests since 1939^{3,8,9}.

Zirconium carbide (ZrC) is a representative non-stoichiometric interstitial compound. It shows high hardness, high melting point, excellent high temperature thermal/mechanical properties, good wear and corrosion resistance, resistance to fission product attack and low neutron cross-section¹⁰⁻¹². It is an important material as high-temperature component and hard coatings^{5,13}, especially a promising candidate as nuclear fuel coating or cladding material¹⁴. Previous studies have found that carbon vacancies significantly affected its mechanical properties^{15,16}, thermo-physical properties¹² and microstructural stability under irradiation¹⁷. As a common phenomenon in MC_{1-x} carbides, the evolution of carbon vacancies dominates the longtime performance and is vital for understanding the high composition deviation.

During the last sixty years, a great number of theoretical and experimental progresses have been paced, in which the stable $V_{\rm C}$ configuration was the key concern. Reviewing the existing literature, we found that the conclusions were typically controversial. For instances, ordering phenomena in $ZrC_{0.51}^{18}$, $ZrC_{0.63}^{19}$, $ZrC_{0.67}^{19}$, $ZrC_{0.74}^{19}$ were claimed to be the same Zr₂C superstructure at first. But later, Obata and Nakazawa proposed that the ordered phase in $ZrC_{0.70-0.75}$ was actually Zr_4C_{32} and there was no Zr_2C ordered phase detected in $ZrC_{0.51}^{20}$. The knowledge was updated recently, *i.e.* the existence of Zr_2C superstructure (*Fd-3m*)⁷ was firmly validated. To understand the mechanism of $V_{\rm C}$ ordering in transition metal carbides, Gusev proposed that the long-range interactions (probably the phonon subsystem) can account for LRO21. Novion held the opposite opinion that VC ordering was dominated by short-range effects³. Up to now, the underlying mechanism pushing forward the evolution of V_{C} configurations is not fully understood.

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Figure 1. Mixing enthalpies (B1-ZrC and *f.c.c.* Zr as reference states) of ordered (the circles) and disordered (the dashed line, $\Delta H_{mix}^{disordered}$) vacancy configurations with various compositions. The ground states are predicted as Zr₈C₇ (*P*4₃32), Zr₆C₅ (*C*2/*m*), Zr₄C₃ (*C*2/*m*), Zr₃C₂ (*Fddd*) and Zr₂C (*Fd-3m*) ordered phases.

In this paper, the energetics of ZrC_{1-x} at various V_C concentrations is studied using the state-of-the-art first principles and cluster expansion method. We report a fingerprint structural unit, namely the V_C triplet constructed by 3rd nearest neighbor carbon vacancies in all the predicted low energy structures of ZrC_{1-x} . The defective structure would lose stability when V_C concentration reaches over 0.5 and simultaneously, highly repulsive V_C s with the 2nd nearest neighbor coordination unavoidably appear. The results explain some longstanding puzzles in non-stoichiometric interstitial compounds, and this study may also shed lights on how to design or tailor the performances of promising transition carbides.

Results

Energetics of ZrC_{1-x}. The mixing enthalpies of diverse ZrC_{1-x} configurations (with reference to B1–ZrC and *f.c.c.* Zr, $0 \le x \le \overline{0.5625}$), as illustrated by circles in Fig. 1, are predicted by cluster expansion method. There are five ground states (GSs), including ordered Zr_8C_7 (P4₃32), Zr_6C_5 (C2/m), Zr_4C_3 (C2/m), Zr_3C_2 (Fddd) and Zr₂C (*Fd-3m*) phases, that restrict the lower bound of mixing enthalpies (see the GS envelope line in Fig. 1, *i.e.* the black curve with circles). These GSs are the configurations with the lowest energy at each composition and will not undergo phase separation into disproportionation products. It would be stated that the ground states of Zr_8C_7 $(P4_{3}2)$, $Zr_{4}C_{3}(C2/m)$ and $Zr_{2}C(Fd-3m)$ are found by exhaustive search in simulation box of $2 \times 2 \times 2$ supercell (32) Zr sites); and $Zr_6C_5(C2/m)$ and $Zr_3C_2(Fddd)$ are disclosed by simulated annealing method in large configuration space with up to 1726 Zr sites ($12 \times 12 \times 12$ supercell). By this way, the predicted GS structure at Zr_6C_5 (C2/m) is isotypic with that of Ti_6C_5 (C2/m)²², and the one at Zr_3C_2 (Fddd) is isotypic with that of Sc_2S_3 (Fddd)²³ (its energy is 4 meV/cation lower than that isotypic with $Ti_3C_2(C2/m)$ predicted in ref. 22). In experiments, only the ordered Zr_2C (*Fd-3m*) phase was characterized by selected area electron diffraction⁷ and neutron diffraction¹⁸ methods. Besides, Obata and Nakazawa observed superlattice lines in annealed $ZrC_{0.7}$ by X-ray diffraction²⁰. They proposed the existence of ordered Zr_4C_3 phase, but did not present the crystal structure. Although predicted Zr_8C_7 (P4₃32), Zr_6C_5 (C2/m) and Zr_3C_2 (Fddd) phases were not found before, their isotypes, V_8C_7 (P4₃32)⁹, Ti_6C_5 (C2/m)²² and Sc_2S_3 (*Fddd*)²³, have been reported.

The energetics of sub-stoichiometric ZrC_{1-x} presents more information for vacancy tolerance and ordering capability. Firstly, ZrC_{1-x} displays significant tolerance to high concentration of V_{CS} . The mixing enthalpy of ZrC_{1-x} with random $V_{\rm C}$ distribution is shown by the dashed curve in Fig. 1. The predicted mixing enthalpies retain negative in the composition range of 0 < x < 0.59. This result suggests that sub-stoichiometric ZrC_{1-x} phases with a huge amount of vacancies are energetically favorable. Otherwise, sub-stoichiometric ZrC_{1-x} with positive mixing enthalpy would spontaneously decompose into f.c.c. Zr and B1-ZrC competition phases. Furthermore, sub-stoichiometric ZrC_{1-x} demonstrates obvious tendency of V_C ordering because numerous V_C configurations have lower mixing enthalpies than the disordered V_C distribution in Fig. 1. The ordering enthalpies of GSs $\left(\Delta H_{ordering}^{GS} = \Delta H_{mix}^{GS} - \Delta H_{mix}^{disordering}\right)$, as represented by the triangles in Fig. 1, are always negative. Meanwhile, with increase of V_C concentration, the ordering enthalpy continuously decreases and reaches the minimum value at around $x \sim 0.5$. This suggests the fact that the higher the V_C concentration, the stronger the ordering tendency of V_{c} s, and in addition, $Zr_{2}C$ has the most obviously ordering tendency in all studied sub-stoichiometric ZrC_{1-x} structures. We expect that ordered Zr₂C should be the most possible ordered phase synthesized in experiments. This result is consistent with the discovery of ordered Zr_2C in experiments^{7,18}. Besides, the calculated ordering enthalpies of Zr_8C_7 and Zr_6C_5 are -30 meV/cation and -55 meV/cation, respectively, which are comparable to that of V_8C_7 and V_6C_5 ordered phases (around -20 meV/cation)³. Because all the GS structures have negative ordering enthalpies, other predicted GS phases would be fabricated by careful controlling of experimental conditions.

Vacancy configurations. Besides the predicted ground states, the structural characteristics of low energy configurations between neighboring GSs are also important to depict the evolution features of V_{CS} in sub-stoichiomitric ZrC_{1-x} . Figure 2 illustrates the radius distribution function of V_C pairs in selected low energy structures with various compositions. V_C configurations are displayed by the presence or absence of certain neighboring V_C pairs,



Figure 2. Radial distribution function of nearest neighboring (NN) V_C pairs in low energy structures (red labels for GSs, black labels for near GSs), as well as the variety of their formation enthalpies. The inset shows the V_C sites in 1NN, 2NN, 3NN and 4NN V_C pairs along {110}, {200}, {211} and {220} crystal directions, respectively.

including 1NN (1st nearest neighboring), 2NN (2nd nearest neighboring), 3NN (3rd nearest neighboring), and 4NN (4th nearest neighboring) V_C pairs. It is striking to notice that 3NN V_C pair is found in all structures, especially only 3NN V_C pair appears in the GS structures of Zr_8C_7 and Zr_6C_5 which have low V_C concentrations. When V_C concentration increases, 1NN V_C pair subsequently presents in the GS of Zr_4C_3 , then 4NN V_C pair emerges in the GSs of Zr_3C_2 and Zr_2C_2 . Besides, 1NN V_C pair is also found in the low energy structures near the GSs, such as $Zr_{32}C_{23}$, $Zr_{32}C_{18}$, and $Zr_{32}C_{17}$. When V_C concentration is higher than 50%, 2NN V_C pair inevitably arises, such as $Zr_{32}C_{15}$ and $Zr_{32}C_{14}$. Accordingly, their formation energies increase sharply with the presence of 2NN V_C coordination.

Figure 2 clearly displays the interesting characteristics of V_C configurations: most frequently observed 3NN configuration, available 1NN and 4NN configurations, as well as the unfavorable or forbidden 2NN configuration. It is important to disclose the global evolution characteristics of V_Cs in binary transition metal carbides. Indeed, it was both experimentally and theoretically found that the vacancies in nonstoichiometric interstitial carbides preferred the 3NN shell and excluded the 2NN shells^{3,22,24}, which was consistent with our calculation results. Hart *et al.* investigated the vacancy distribution in low energy structures of TiC_{1-x} using cluster expansion method. Although they did not mention the evolution of V_C configurations at various concentrations, carbon vacancies were found arranging themselves in [112] rows²⁵, which was along the crystal direction of 3NN V_C pairs, and 2NN configurations gradually disappeared during annealing or ordering.

We expect that there would be inherent local V_C configurations commonly appearing in the low energy structures because their mixing enthalpies are so close to the GS envelope line. In ZrC_{1-x} with stoichiometry near ordered GS phases (around compositions of Zr_2C , Zr_4C_3 , Zr_3C_2 etc.), it would be unusual for any abrupt change of V_C patterns. After careful analysis of V_C configurations, the common configuration unit, *i.e.* the 3NN V_C triplet, is identified. Figure 3 illustrates the overall features of V_C distributions in C-sublattice. In Zr_8C_7 ($P4_332$) and Zr_6C_5 (C2/m) as shown in Fig. 3(a,b), respectively, neighboring V_C s only occupy 3NN sites and form the 3NN V_C triplets. The schematic of 3NN V_C triplet is displayed in Fig. 3(f), which is an equilateral triangle with its side length restricted by 3NN coordination. Although it was known that the preference of occupying 3NN V_C shell was typical in ordered structures of sub-stoichiometric transition metal carbides and nitrides, the significance of V_C triplet configuration was not reported before. We find that the 3NN V_C triplet prevails in all the low energy structures like a fingerprint V_C configuration.

The 3NN V_C triplets are corner-shared in Zr₈C₇ (*P*4₃32) (as shown in Fig. 3(a)), and edge-shared in Zr₆C₅ (*C2/m*) (as shown in Fig. 3(b)). In Zr₆C₅ (*C2/m*), one defective carbon layer (filled with 2/3 carbon atoms) and its neighboring perfect carbon layer stack alternatively along $< 211 > _{B1}$ crystal direction. When V_C concentration is higher than 1/6, V_Cs could not be accommodated by occupying the sites restricted by pure 3NN coordination. Neighboring 3NN V_C triplets adjust their relative positions and bring out the 1NN V_C configuration to host more V_Cs, as shown for Zr₄C₃ (*C2/m*) in Fig. 3(c). The 3NN V_C triplets are predominant in the {111}_{B1} carbon layers and stack along $< 211 > _{B1}$ crystal direction. With more V_Cs incorporated in Zr₃C₂ (*Fddd*), 3NN V_C triplets link adjacent defective carbon layers, as shown in Fig. 3(d). Additionally, self-assembling of 3NN V_C triplets generates the 1NN and 4NN V_C configurations. For Zr₂C (*Fd-3m*) shown in Fig. 3(e), neighboring carbon layers have 1/3 and 2/3 C-sublattice sites occupied by V_Cs, respectively, in which crowded 3NN V_C triplets are orthocenter overlapped. It is noted that every carbon atom in Zr₂C (*Fd-3m*) is surrounded by overlapped 3NN V_C triplets. These 3NN V_C triplets are completely coordinated by 1NN and 4NN V_C configurations. Changing any V_C site or forming one more V_C will introduce the 2NN coordination. Besides these GS structures, remarkable 3NN V_C triplets are also found as the common configuration unit in near GS configurations, such as Zr₃₂C₂(*C2*) and Zr₃₂C₂₃(*R-3m*).

Short-range interactions. The above results illustrate the self-assembling of V_C configuration and feature the fingerprint configuration of 3NN V_C triplet in defective ZrC_{1-x} . It is speculated that the redistribution of electrons



Figure 3. Vacancy configurations in C-sublattice of (a) Zr_8C_7 (*P4*₃32), (b) Zr_6C_5 (*C2/m*), (c) Zr_4C_3 (*C2/m*), (d) Zr_3C_2 (*Fddd*) and (e) Zr_2C (*Fd-3m*), with significant character of (f) 3NN V_C triplet.



Figure 4. Interaction energies of 1NN, 2NN, 3NN, 4NN V_C pairs (the circles) and 3NN V_C triplet (the star); the inset shows that neighboring Zr atoms of 3NN V_C triplet are endowed with the square-pyramid coordination (C_{4v}).

around vacancies^{22,26,27} would affect the short-range interactions among V_Cs by altering the M-C bonds, which would be the important driven force pushing forward the local ordering pattern of V_Cs. Therefore, the interaction (or binding) energies of various V_C clusters, and their correlations with the evolution of V_C configurations, are studied here. Using the $3 \times 3 \times 3$ supercell with 108 Zr sites, the interaction energies of V_C pair and triplet are calculated by the following equations²⁸:

$$E_{i}(V_{C} \text{ pair}) = [E(Zr_{108}C_{108}) + E(Zr_{108}C_{106}) - 2 \times E(Zr_{108}C_{107})]/2$$
(1)

$$E_{i}(V_{C} \text{ triplet}) = [2 \times E(Zr_{108}C_{108}) + E(Zr_{108}C_{105}) - 3 \times E(Zr_{108}C_{107})]/3$$
(2)

where the first and last terms on the right side are the total energies of perfect ZrC and the supercell containing an isolated V_c , respectively; and the second items in equations (1) and (2) stand for the total energies of the supercells containing various V_c pairs and the 3NN V_c triplet, respectively. The derived interaction energy indicates the thermal stability of various vacancy clusters relative to isolated V_c s with infinite distance. and determines whether isolated vacancies would aggregate together to form certain vacancy configuration.

Figure 4 plots the interaction energies of various V_C pairs and the 3NN V_C triplet. For V_C pairs, interaction energies increase in the order of $E_i(3NN) < 0 < E_i(1NN) < E_i(4NN) < E_i(2NN)$, which agrees with that reported by Razumovskiy *et al.*^{29,30}. The interaction energy of 3NN V_C triplet yields -49 meV/vacancy or -147 meV/triplet, which is lower than that of 3NN V_C pair (-22 meV/vacancy or -44 meV/pair). Therefore, the formation of 3NN V_C triplet benefits to lower the energy of defective structure. The inset in Fig. 4 sketches neighboring Zr_6C



Figure 5. The projected density of states (PDOS, unit states/eV·atom) and the projected crystal orbital Hamilton population (pCOHP) of nearest-neighbor Zr-C interactions in (**a**) ZrC (*Fm-3m*), (**b**) Zr₂C (*Fd-3m*) and (**c**) Zr₃₂C₁₅ (*R-3m*). The solid lines in pCOHP diagram stand for the sum of *pd* bands, while the shading zones denote the contributions from certain orbitals (No scale is given for the pCOHP analysis).

octahedra with a 3NN V_C triplet. The C atoms in octahedra are removed to produce V_Cs, and the 3NN V_C triplet configuration ensures each Zr atom coordinated with only one V_C. Therefore, the carbon coordination of Zr atom is minimally disturbed, and thereby the bonding feature is least affected. Meanwhile, the self-assembling of V_Cs allows short-range ordering by coordinating the most preferred V_C configuration, *i.e.* 3NN V_C triplet. Local ordering in the configuration of 3NN V_C triplets would be universal in ZrC_{1-x} at any composition. In elastic diffuse neutron scattering experiments, $ZrC_{0.80}$ and $ZrC_{0.64}$ were identified with the same peak positions although they have different compositions³. This result suggests similar short-range ordering in the two compounds. If the short-range ordering covers the whole lattice and the defective structure satisfies a new symmetry at certain composition, then an ordered ZrC_{1-x} phase would be identified. All the results show that self-assembling of 3NN V_C triplets is the key factor to maintain phase stability of defective ZrC_{1-x} with high concentration of V_Cs, and to realize short-range ordering or dering of V_Cs.

The interaction energy of 1NN V_C pair, 13 meV/vacancy, is slightly higher than zero but much lower than those of the 4NN and 2NN V_C pairs, which are 70 meV/vacancy and 321 meV/vacancy, respectively. The small value of 1NN V_C pair shows very weakly repulsive interaction. Therefore, 1NN V_C pair is the second preferred configuration in the low energy structures. With more V_Cs accommodated in GS structures of Zr₃C₂ and Zr₂C, together with near GS structures Zr₃₂C₂₃, Zr₃₂C₁₈ and Zr₃₂C₁₇, both 1NN and 4NN V_C pairs present in order to efficiently coordinate neighboring 3NN V_C triplets. Therefore, the presences of 1NN and 4NN V_C configurations are helpful for balancing V_C concentration and thermal fluctuation.

The 2NN $V_{\rm C}$ pair has extremely high interaction energy (321 meV/vacancy), which means significantly repulsive interaction between $V_{\rm C}s$ in 2NN configuration. Therefore, 2NN $V_{\rm C}$ pair is unfavorable or forbidden in defective ${\rm ZrC}_{1-x}$ and it does not show up in low energy structures when $V_{\rm C}$ concentration is lower than 50%. It's noteworthy that the strongly repulsive interaction among 2NN configurations may prevent the formation of large-scale $V_{\rm C}$ clusters in sub-stoichiometric ${\rm ZrC}_{1-x}$. Unfortunately, 2NN coordination is inevitable when $V_{\rm C}$ concentration is higher than 50%. The appearance of 2NN configuration will abruptly increase the formation enthalpy (shown in Fig. 2) and therefore, the defective ${\rm ZrC}_{1-x}$ may undergo phase separation in such case. This might be the reason why the critical $V_{\rm C}$ concentration is limited around 50% in ${\rm ZrC}_{1-x}$.

The short-range interactions among vacancies could well explain the evolution feature of V_C configurations and the fingerprint configuration. The result indicates that these are the driven-force of V_C self-assembling, *i.e.* bringing down the system energy *via* the maximization of attractive 3NN configuration and the exclusion of strongly repulsive 2NN coordination, and simultaneously balancing 3NN V_C triplets through the moderately repulsive 1NN and 4NN interactions. Also, the short-range interactions among V_C s would be more and more significant at high V_C concentration. It may provide hints on the origin of enhanced ordering tendency with increasing V_C s.

Electronic structures of ZrC_{1-x}. Projected density of states (PDOS) and projected crystal orbital Hamilton population $(pCOHP)^{31}$ are illustrated in Fig. 5 to describe the Zr-C bonding characters in highly defective ZrC_{1-x} . At high V_C concentration, the electronic structures of ZrC_{1-x} would be significantly affected by vacancy-vacancy interaction and/or vacancy ordering. This was verified by Pickett and Klein as they found complex differences between the calculated electronic structure of an isolated carbon vacancy in B1-NbC by the muffin-tin Green's-function method and the experimental electronic structure of NbC_{0.85} from X-ray photoemission spectrum³². Therefore, we investigate the electronic structures of ordered Zr_2C and $Zr_{16}C_{15}$ phases to understand the

electronic structures of ZrC_{1-x} around the critical composition of x = 0.5. This would illustrate the mechanism of electronic redistribution at high V_C concentration.

Firstly, we study the bonding characters in perfect ZrC (B1) as shown in Fig. 5(a). In the perfect unit cell, Zr atom is octahedral coordinated by six C atoms with a high site symmetry (O_h), which ensures good capability of band overlapping. The bonding region in PDOS shows a strong hybridization of C *p*- and Zr *d*-states, which promotes the formation of strong covalent Zr-C bond. Besides, the orbital-wise pCOHP analysis indicates that Zr-C bond is dominated by the *p*-*d* σ and π bonding with significant share of C(*p*_z) and Zr(*d*_{3z²-r²}) interactions. These bonding characters are consistent with other transition metal carbides like TiC^{26, 33}.

With removing of C atoms, *i.e.* the electron acceptors in ZrC_{1-x} , the excess *d* electrons redistribute on electronic states in high energy level, as clearly shown in Fig. 5(b). For Zr_2C at the critical concentration, the bonding states between $-2.4 \, \text{eV}$ and $-4.8 \, \text{eV}$ are mainly dominated by $\operatorname{C}(p_z)$ and $\operatorname{Zr}(d_{3z^2-r^2})$, and the PDOSs locating from $-1.9 \, \text{eV}$ to Fermi level (E_F) correspond to the *d*-*d* bonding among Zr atoms. When the concentration of V_Cs is higher than 50%, for example $\operatorname{Zr}_{32}C_{15}(R-3m)$ with inevitable 2NN V_C coordination, anti-bonding states originated from $\operatorname{C}(p_z)$ - $\operatorname{Zr}(d_{xz})$ interactions obviously show up near E_F in Fig. 5(c). As a result, the formation enthalpy is quite high for ZrC_{1-x} with x > 0.5, which contains unfavorable 2NN V_C configuration.

Discussion

We found neighboring V_{C} s have totally different values of short-range interactions, which increase in the order of $E_i(3NN) < 0 < E_i(4NN) < < E_i(2NN)$. It goes along with the evolution characteristics of V_C configurations. The moderately attractive and strongly repulsive interactions between V_C s in 3NN and 2NN configurations, respectively, provide the driven-force of self-assembling of V_C s. Meanwhile, 3NN V_C triplet is more stable than other V_C configurations, and it serves as the fingerprint block in low energy ZrC_{1-x} structures. At high V_C concentration, neighboring 3NN V_C triplets modify relative positions by coordinating 1NN and 4NN V_C configurations. The present results clearly demonstrate that ordering of V_C s in ZrC_{1-x} is not an abrupt structural change. The local ordered configurations, long-range ordering spreads throughout the whole lattice and one could observe ordered phase with new space group. The underlying mechanism falls into the concept of self-assembling of V_C s. Short-range interactions among V_C s are the driven-force of local or long-range ordering; and the formation enthalpy is reduced *via* the generation of attractive 3NN interactions and the exclusion of strongly repulsive 2NN interactions.

It is crucially important to realize the local ordering of $V_{C}s$ and to maintain phase stability in highly sub-stoichiometric ZrC_{1-x} . On one hand, local ordering of V_{C} configuration with 3NN triplets, instead of the disordered distribution, stabilizes the defective structures by minimally affecting bonding features. On the other hand, the strongly repulsive interaction between $V_{C}s$ in 2NN configuration weakens the formation of large scale vacancy clusters or voids as well. More encouragingly, the self-assembling of $V_{C}s$ provides us the opportunity to tailor overall properties through defect engineering. One exciting perspective is the improvement of radiation resistance of ZrC_{1-x} via the mechanism of vacancy mediated performance optimization. Local ordering of $V_{C}s$ benefits the accommodation, annihilation and recombination of radiation-induced C-related point defects. Therefore, ZrC_{1-x} with optimal composition may be promising for next generation nuclear fuel coating and cladding material. Another important prospect is to precisely tune mechanical and thermal properties through adjusting the chemical composition of ZrC_{1-x} . In the near future, it demands extensive investigations on the effects of V_{C} concentrations and configurations on radiation resistance and thermo-physical properties.

We herein proposed the physical insight of the critical V_C concentration, $x \sim 0.5$, in ZrC_{1-x} . At relatively low V_C concentration, excess *d* electrons redistribute on *d-d* bonding states after the removal of C atoms; whilst neighboring Zr-C bonding is not affected significantly. This mechanism is helpful to maintain the structural stability of defective ZrC_{1-x} . When the V_C concentration is higher than the critical composition, 2NN V_C configuration inevitably appears which brings out anti-bonding states between Zr and C atomic interactions. The anti-bonding states lead to extremely high formation enthalpy of defective ZrC_{1-x} with 2NN V_C coordination and thereby, the ZrC_{1-x} compound loses stability and undergoes phase separation.

The current investigation suggests new clue for better understanding of the phase stability and defect engineering in non-stoichiometric interstitial compounds. Firstly, the phase stability and ordering mechanism in ZrC_{1-x} , especially the discovery of self-assembling of 3NN triplets driven by short-range interactions among $V_{C}s$, may be common in binary transition metal carbides. There are solid evidences on the similar V_C configurations in non-stoichiometric carbides, for instances, the isotopic structures of V_8C_7 ($P4_332$)⁹ and Ti_6C_5 (C2/m)²² with Zr_8C_7 ($P4_332$) and Zr_6C_5 (C2/m), respectively; the widespread occupation of 1NN and 3NN shells, but not 2NN shells in TiC_{1-x} and ZrC_{1-x}^6 ; as well as the same neutron diffuse scattering in $TiC_{0.5-0.6}$, $VC_{0.75}$ and $NbC_{1-x}^{-3.34}$. This work presents hints to understand following puzzles, like why common characteristics of vacancy configurations exist and what are mechanisms of phase stability and V_C ordering in sub-stoichiometric binary transition metal carbides. Secondly, it needs thorough inquiry whether the present results could be extended to other sub-stoichiometric binary transition metal nitrides and oxides because short-range interactions among anion vacancies are possibly different. Various self-assembling mechanisms of anion vacancies may dominate the diversity of the ordering phenomena in sub-stiochiometric nitrides and oxides.

In summary, vacancy configurations and their evolution in sub-stoichiometric ZrC_{1-x} were investigated by combining first-principles calculations, cluster expansion and supercell methods. Firstly, the negative mixing enthalpy and the negative ordering enthalpy are direct energetic proofs of the tolerance and the ordering tendency of $V_C s$ in sub-stoichiometric ZrC_{1-x} . Besides, the energetically preferred 3NN V_C triplet is the fingerprint structural unit in sub-stoichiometric ZrC_{1-x} . To balance V_C concentration and thermal fluctuation, 1NN and 4NN V_C configurations with moderately repulsive interactions would show up. The tolerance of vacancies is limited at around 50% because the presence of unfavorable 2NN V_C coordination with strongly repulsive interaction energy would lead to structural instability or phase separation.

It's noteworthy that the self-assembling of 3NN V_C triplets driven by short-range interactions is the fundamental essence of the phase stability, short-range and long-range ordering in sub-stoichiometric ZrC_{1-x} . For the first time, we disclose the significant self-assembling mechanism of V_Cs in sub-stoichiometric ZrC_{1-x} which provides us the opportunity to tailor its properties, such as the radiation resistance, mechanical and thermal properties, through defect engineering.

Methods

Sub-stoichiometric ZrC_{1-x} is taken as a pseudo "binary solid solution" consisting fully occupied Zr-sublattice and mixing of carbon atoms and vacancies at C-sublattice. This system can be handled by supercell^{4,35}, order parameter functional method (OPF)³⁶, coherent-potential approximation³⁷, or cluster expansion (CE)³⁸ methods. Three ordered phases, Zr₂C, Zr₃C₂ and Zr₆C₅, have been predicted using OPF method³⁶, but the results did not provide structural information. Theoretical calculations using supercell method^{4,35} has emphasized the importance of lattice relaxation effect and the vacancy distribution, however it may leave out important configurations because of limited trial structures. The CE method which is the choice here, considers the effects from short-range interactions and local relaxation, and therefore provides both the structural characteristics and corresponding structural energy with DFT (Density Functional Theory) accuracy³⁹.

Within the CE approach, the configuration on C-sublattice is described by configurational pseudospin variable σ_i for lattice site *i*, which takes value +1 or -1 depending on the occupation of lattice site *i* by carbon atom or carbon vacancy. A particular arrangement of spins is called a configuration (or structure), which is represented by a vector $\{\sigma_i\}$ containing the value of configurational pseudospin variable for each site *i*. The energy of a given composition and configuration is expressed as^{39,40}:

$$H = V_0 + \sum_i V_i \sigma_i + \sum_{i,j} V_{ij} \sigma_i \sigma_j + \sum_{i,j,k} V_{ijk} \sigma_i \sigma_j \sigma_k + \dots = \sum_{\alpha} V_{\alpha} \sigma$$
(3)

where V_{α} and φ_{α} are called the effective cluster interaction (ECIs) and the correlation function of cluster α (corresponding to a set of sites *i*: pairs, triplets, tetrahedral, *etc.*), respectively. For disordered phases, the energy is only composition (*x*) dependent⁴¹:

$$H = \sum_{\alpha_n} V_{\alpha_n} (1 - 2x)^n \tag{4}$$

where α_n stands for one kind of n-body cluster.

A principal objective of the CE method is to evaluate unknown ECIs so that the configuration characters can be predicted with the accuracy of DFT calculations. Here, we determine the ECIs in the framework of the structure inversion method⁴²⁻⁴⁴. When energies of ordered structures are prepared from DFT calculations, the ECIs are determined by fitting a truncated form of equation (3) to the DFT energies using the least-squares technique. The accuracy of CE prediction is assessed by the leave-one-out and leave-many-out cross-validation (CV) score⁴²⁻⁴⁴. The set of clusters is optimized using genetic algorithm *via* minimizing the CV score. The DFT structures include initial selected random structures, low energy structures refined during construction of CE, and reported superstructures in non-stoichiometric interstitial compounds^{9,45}. Finally, 25 clusters and 250 structures were selected to construct the CE formula. The average prediction error of the final CE is less than 4.3 meV/cation (per Zr/C-sublattice site). Thereafter, two types of calculations are used to search low energy structures (LESs) *via* the constructed CE. One is an exhaustive search by calculating the energies of all possible configurations in a finite-sized cell. The other is the simulation annealing (SA) method⁴⁶ within a larger structural space. Here, exhaustive search was used for the supercell containing up to 1728 Zr sites. Detailed procedure of constructing CE can be found in ref. 47. In this work, CLUPAN code^{42,48} is used for the construction of CE and the searches of LESs.

First-principles calculations were done using VASP⁴⁹ code, in which the projector augmented-wave (PAW) method^{50,51} within generalized gradient approximation⁵² was employed. The plane-wave basis cutoff and the *k*-mesh separation were set as 500 eV and 0.04 Å⁻¹, respectively. Full structural relaxations (atomic positions and lattice constants) were performed until the energy difference converges to less than 10^{-6} eV.

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Author Contributions

J.W. conceived the project. Y.Z. performed all calculations. B.L. advised CE calculation. Y.Z., B.L. and J.W. analyzed the data, discussed the results and wrote the paper.

Additional Information

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