How does conformational flexibility influence key structural features involved in activation of anaplastic lymphoma kinase?†

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Anaplastic Lymphoma Kinase (ALK) plays a major role in developing tumor processes and therefore has emerged as a validated therapeutic target. Applying atomistic molecular dynamics simulations on the wild type enzyme and the nine most frequently occurring and clinically important activation mutants we revealed important conformational effects on key interactions responsible for the activation of the enzyme.

Introduction

Anaplastic Lymphoma Kinase (ALK) is a member of the insulin receptor tyrosine kinase superfamily that normally participates in the development of the nervous system in embryos, the expression of which decreases after birth.1,2 Twenty different ALK-fusion proteins resulting from different chromosomal rearrangements have been shown to be connected with the pathogenesis of diseases such as anaplastic large-cell lymphoma, diffuse large B-cell lymphoma and inflammatory myofibroblastic tumors.3 A series of germline and somatic single nucleotide polymorphisms in the catalytic domain of ALK were characterized recently from neuroblastoma patient samples and many were shown to be related with deregulation of ALK and its constitutive activation.4–6 Recently resolved crystal structures of the wild type enzyme,7 its complexes with adenosine diphosphate (ADP), and staurosporine (STU),7 adenosine triphosphate (ATP) competitive inhibitors – PHA-E429, NVP-TAE6848 and of two activation mutants9 revealed a canonical protein kinase domain fold and topology. The molecule is built of a smaller N-terminal lobe and a larger C-terminal lobe (Fig. 1).7 The N-terminal lobe contains five stranded antiparallel β-sheets, an important regulatory αC-helix which can be in active and inactive positions and an ATP-binding glycine-rich loop (P-loop). The juxtamembrane segment of the N-terminus forms a β-turn motif which is unique for ALK. The C-terminal lobe contains eight α-helices (αD–αI) and two β-stranded sheets. The activating loop (A-loop) which contains residues 1270–1299 is located here and starts with a DFG-motif, followed by a short αA-helix and ends with a Pro-Pro-Glu motif. The catalytic loop is located between αE and the first strand of the β-sheet in the C-lobe. The αA-helix is located below and orthogonally to the αC-helix and against the N-terminal β-turn by multiple polar and hydrophobic interactions.

The crystal structures display several features of inactive kinase conformation such as the degree of lobe closure between N- and...
C-terminal lobes, the position of the αC helix and its orientation with respect to the C-lobe and the location of the distal part of the A-loop which obstructs the peptide binding site. The structure does not possess, however, all the negative regulatory elements that exist in the insulin receptor kinases. The A-loop is in an intermediate position between the inactive and active kinases and its proximal part does not block the ATP-binding site as in other inactive kinase conformations. The DFG-motif exhibits a “DFG-in” conformation as in the case in the active kinases, and the αC-helix allows the proper positioning of Glu1167 with Lys1150 for salt-bridge formation, which is also typical for the active conformation.1

ALK activation involves complex changes in several structural features and interactions including but not limited to the mutual orientation of the αC and αA helices, the position of the A-loop, the hydrogen bond between Tyr1278 and Cys1097 from the N-terminal β-turn which locks the inactive conformation and the salt bridge between Glu1167 and Lys1150 which is distinctive for the active conformations. Lack of insights into conformational dynamics is a major limitation of X-ray protein crystal structures as they are largely static in nature and reflect some of the details of the experimental methodology, such as ensemble and time averaged properties, imposition of crystal packing effects, and some deviations due to the non-linearity of the registered signals and methods of interpretation. Most importantly, crystal structures could miss some important information about the conformational dynamics and relaxation which are fundamental protein properties. The delicate interactions which define such unique intermediate conformations would be influenced by the internal conformational flexibility under native conditions; therefore exploration using molecular dynamics (MD) is important in order to complement the crystallographic structural information. In addition the correlation between the protein dynamics in enzyme catalysis is matter of crystallographic structural information. In the case in the active kinases, and the αC-helix allows the proper positioning of Glu1167 with Lys1150 for salt-bridge formation, which is also typical for the active conformation.7

The focus of our study is to establish whether the structure of ALK, revealed by crystallographic studies as an intermediate between the active and inactive conformations, is altered by the conformational flexibility and to define the interactions responsible for this. To achieve this insight we carried out a comprehensive set of atomistic molecular dynamics (MD) simulations followed by detailed analysis of the time evolution and flexibility of multiple interactions representative for both the inactive and active ALK conformations. Here we discuss the influence of the conformational flexibility of the key structural features of the inactive and active conformations whilst the more comprehensive structure/mobility/to activity relationship would be an objective of further project.

Methods

The MD simulations were carried out with the GROMACS code (version 4.3.1)18 and Gromos43b1 force field for 100 nanoseconds (ns). Twelve MD simulations were carried out for the wild-type apoenzyme ALK, ALK–ADP complex, ALK–staurosporine complex and nine activation mutations – M1166R, I1171N, F1174L, F1245C, I1250T, G1128A, T1151M, R1192P and R1275Q.4–6 The system was prepared from the following crystal structures of ALK taken from PDB19 as follows: for the wild type free ALK MD simulation 3L9P.pdb structure was used; for the ALK–ADP complex the 3LCT.pdb structure which contains the ligand ADP was used; for ALK–staurosporine complex, 3LCS.pdb structure which contains the inhibitor staurosporine was used; for the nine simulations of the activation mutant forms of the free ALK the 3L9P.pdb structure was used from which all nine structures of activation mutations were prepared from the crystal structure of the wild-type enzyme using the “What if” web interface (http://swift.cmbi.ru.nl/servers/html/index.html).20 In all simulations the missing residues in the crystal structures of ALK were added using PyMol.21 Gromacs utilities for system preparation were used thoroughly. The hydrogen atoms were added according the protonation states of the ionogenic groups and consequently minimized. Gromos43a1 force field was thoroughly applied. The ADP and staurosporine parameters were generated using PRODRG web-based tool.22 Subsequently the entire system was minimized using the steepest descent algorithm. The system was then solvated using rectangular SPC23 water box placed 10 Å from the edges of the protein and neutralized. The solvent was then equilibrated for 50 picoseconds (ps) at 310 K using NVT ensemble. Production MD simulations were run for 100 ns in NPT ensemble at 310 K applying Berendsen thermostat24 treating the electrostatic interactions by particle mesh Ewald method. The simulation step was 2 fs and nonbonding cutoff of 14 Å was applied. The simulation system is presented on Fig. 2.

The quality of the simulations was assessed by Root Mean Square Deviation (RMSD) (Fig. S1–S3A–I, ESL†), root mean
square fluctuation (Fig. 3 in the main text and Fig. S4A–D and S5A–D in ESI†), radius of gyration and total, kinetic and potential energy values long the time of the simulation (the plots are available from the authors by request). All calculations were done for the αC atoms in reference to the initial minimized structure. The stability of hydrogen bonds during the simulations was assessed using Gromacs g_hbond utility. After that the average number of the hydrogen bonds between particular couples of residues per MD trajectory was extracted. Salt-bridge distances were calculated using g_salt utility and the averaged value for a particular couple of residues was calculated for each trajectory. Several hydrophobic interactions were calculated by g_dist command. The cluster analysis was done using cutoff distance of 0.25 nm and the central structure of the largest cluster was used for comparison with the minimized crystal structure.

Results and discussion

The N-terminal β-turn, followed by the glycine-rich (P-loop) and the A-loop (Fig. 3) exhibit high flexibility in the wild type free enzyme (based on the RMSF values) which suggest that during the process of activation they would undergo sensitive changes. In the simulations of the structures with activation mutants the N-terminal β-turn is characterized with higher flexibility than the wild type enzyme in all mutants with the exception of I1171N and T1151M (Fig. S5 in ESI†). The flexibility of the A-loop is also increased by the activating mutations (with exception of G1128A). The P-loop is characterized with increased flexibility in five mutants (I1250T, R1275Q, M1166R, R1192P and T1151M). The N-terminal β-turn, glycine-rich (P-loop) and the A-loop which are very flexible in the wild type enzyme and as a result of the activating mutations their flexibility is increased even more. The binding of ADP and staurosporine does not influence the flexibility of the catalytic loop, αA- and αC-helices (Fig. S4 in the ESI†) but (especially the binding of staurosporine) leads to increased flexibility of the P-loop and the A-loop (excluding the αA-helix).

The orientation between the αC- and αA-helices which is in is key importance for switching between the inactive and active conformations is stabilized by hydrogen bonds and electrostatic interactions (Fig. 4). An insight about the orientation of the αA and αC helices is provided by analysis of the hydrogen bonding and electrostatic interactions between them (Tables 1 and 2). The hydrogen bond between the side chain of Arg1275 and the carbonyl oxygen of Asp1163 is presented in 60% and 70% from the MD trajectories of T1151M and R1275Q mutants, 40% in the wild type enzyme and the F1245C, I1250T, I1171N mutants and with lower frequency in the G1128A, M1166R and F1174L. Arg1279 interacts with Asp1163, as in the crystal structure, which is indicated by both the hydrogen bonding profile and the salt-bridged distance but this interaction tends to become less strong. In the wild type Arg1275 and Glu1167 still interact electrostatically and by a hydrogen bond, however in most of the mutant forms, this hydrogen bond is present with lower frequency. The hydrogen bond between Arg1279 and Gln1159 is stable in the MD trajectory of the wild type ALK and is presented with differential stability in the mutants (e.g. it is relatively stable in FF1174L and T1151M but is missing in G1128A and R1275Q).

The side chain of Asp1276 rotates towards the C-lobe during the simulation and the hydrogen bond between Arg1275 and Asp1276, which exists in the crystal structure, is not presented in the simulation of the wild type and mutant forms (the averaged distance between the side chains is higher than in the crystal structure; 4.2 Å compared to 2.0 Å for the wild type and even large differences in the mutant forms). The hydrogen bond between Lys1285 from the A-loop and the αC-helix residue Asp1163, disappears during the simulation of the wild type and all activation mutant enzyme forms and Lys1285 moves away from the αC-helix, due to the flexibility of the A-loop. The repulsive interactions between the side chains of Arg1275 and
Arg1279 with the side chain of Lys1285 might also contribute to this effect. The balance between these repulsive interactions and the attractive interactions between the side chains of Lys1285 and Asp1163, Asp1160, Glu1167 and Asp1276, together with the high flexibility of the A-loop, can be a controlling instrument for the positioning of the distal part of the A-loop in more active-like orientation and thus postulated to be part from the activation mechanism. The hydrogen bonding and electrostatic interactions between \( \gamma \)C and \( \gamma \)A helices become less strong and many of them partly or almost disappear during the simulations of the wild type and mutant forms. This could be an initial step from more long-range process of reorientation of the \( \gamma \)C and \( \gamma \)A helices as a part from the enzyme activation.

An important interaction that locks the \( \gamma \)C helix in inactive conformation is the hydrogen bond between Tyr1278 from the A-loop (the preferred tyrosine residue for the autophosphorylation reaction), and Cys1097, from the N-terminal \( \beta \)-turn and the central structure of the most populated cluster (Fig. 5). While stable during 60% of the wild type ALK MD simulation and is even much stronger in all mutant forms than in the crystal structure during the simulation of the wild type enzyme and even much stronger in all mutant forms (Table 1) which indicate that the structure became more stable during 60% of the wild type ALK MD simulation and is even much stronger in all mutant forms than in the crystal structure during the simulation of the wild type enzyme and even much stronger in all mutant forms (Table 1) which indicate that the structure became more stable during 60% of the wild type ALK MD simulation and is even much stronger in all mutant forms than in the crystal structure during the simulation of the wild type enzyme and even much stronger in all mutant forms than in the crystal structure during the simulation of the wild type enzyme. The existence of this interaction in ALK is in contrast to all known active kinases.

Importantly, the hydrogen bond between the second A-loop tyrosine Tyr1282 and Arg1284 also almost disappears in the simulations of the wild type and the mutant forms of the enzyme which also might be part from the initiation events of the activation. This could be facilitated by the flexibility of the A-loop and the stronger interactions between the side chains of Arg1284 and Glu1303 and between the side chains of Tyr1282 and Arg1248. In contrast the hydrogen bond, representative for the inactive conformation between the peptide backbones of DFG + 1 residue Met1273 and Ile1277 that exists in the crystal structure is preserved over the simulation with very high frequency and is also present in all activation mutants. The existence of this interaction in ALK is in contrast to all known active kinases.

Distinctive interaction for the active kinases conformations is the salt bridge between Lys1150 and Glu1167. It is stronger than in the crystal structure during the simulation of the wild type enzyme and even much stronger in all mutant forms (Table 1) which indicate that the structure became more active-like regarding this interaction.

The comparison between the minimized crystal structure and the central structure of the most populated cluster (Fig. 5)
from the simulation of the wild type enzyme confirms that the A-loop moved more towards the active position, which is more expressed in its distal part during the simulation, and the P-loop moved more towards the catalytic loop, whilst the inter-lobe did not change considerably during the simulation. Two interactions might be related to the positioning of the P-loop closer to the catalytic loop. The salt bridge between Lys1150 and Glu1167, which is characteristic for the active conformation, is strengthened during the MD simulation of the wild type and all mutants. In addition Arg1120 switches between two glutamate residues – Glu1132 and Glu1129.

The results provide important insight revealing that numerous interactions undergo initial changes relevant to the activation, namely: the hydrogen bonds and electrostatic interactions which keep the inactive orientation between αC and αA helices are less strong than in the crystal structure as well as the hydrogen bond between the αC-helix to the N-terminal β-turn; the A-loop is displaced in the more active position due to the repulsive interactions between Lys1150 and Arg1279. Essentially, the salt-bridge between Glu1167 and Lys1150 which is characteristic for the active conformation is strengthened and might help for displacing of the P-loop towards the catalytic loop. The N-terminal β-turn which is unique for ALK exhibits high flexibility in the wild type ALK and all mutant forms. The activating mutations tend to increase even more the conformational flexibility with most profound effect of the A-loop and N-terminal β-turn and the P-loop confirming their role during the activation process. The complex process of activation might be initiated by several events such as loosening the electrostatic interactions and hydrogen bonding between the αC- and αA-helices, increasing the flexibility of the A-loop and displacing it in a more active position; increasing the flexibility of the N-terminal β-turn in combination with the loosening and breaking of the “inactive conformation” hydrogen bond between Tyr1278 and Cys1097, strengthening the “active conformation interaction” between Glu1167 and Lys1150; and displacing of the P-loop towards the active site. The gained atomistic insight can be used for further atomistic and mechanistic studies and as a background in designing more effective ALK inhibitors as potential drugs.

CC and TK would like to thank HPC-Europe Program, UK National Supercomputer Hector, UK National Service for Computational Chemistry Software, their Marie Curie International Outgoing Career Development Fellowships and the HPC-cluster at the Department of Applied Sciences at Northumbria University at Newcastle. CC would like to acknowledge Fulbright Senior Research Grant from Bulgarian-American Fulbright Commission.

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