Structural communication between the GTPase Sec4p and its activator Sec2p: Determinants of GEF activity and early deformations to nucleotide release

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Abstract

Ras GTPases are molecular switches that cycle between OFF and ON states depending on the bound nucleotide (i.e. GDP-bound and GTP-bound, respectively).

The Rab GTPase, Sec4p, plays regulatory roles in multiple steps of intracellular vesicle trafficking. Nucleotide release is catalyzed by the Guanine Nucleotide Exchange Factor (GEF) Sec2p. Here, the integration of structural information with molecular dynamics (MD) simulations addressed a number of questions concerning the intrinsic and stimulated dynamics of Sec2p and Sec4p as well as the chain of structural deformations leading to GEF-assisted activation of the Rab GTPase.

Sec2p holds an intrinsic ability to adopt the conformation found in the crystallographic complexes with Sec4p, thus suggesting that the latter selects and shifts the conformational equilibrium towards a pre-existing bound-like conformation of Sec2p.

The anchoring of Sec4p to a suitable conformation of Sec2p favors the Sec2p-assisted pulling on itself of the a1/switch I (SWI) loop and of SWI, which loose any contact with GDP. Those deformations of Sec4p would occur earlier. Formation of the final Sec2p-Sec4p hydrophobic interface, accomplishes later.

Disruption of the nucleotide cage would cause firstly loss of interactions with the guanine ring and secondly loss of interactions with the phosphates. The ease in sampling the energy landscape and adopting a bound-like conformation likely favors the catalyzing ability of GEFs for Ras GTPases.

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1. Introduction

Ras GTPases or G proteins (Guanine Nucleotide-Binding Proteins (GNBP)) are molecular switches that cycle between OFF and ON states depending on the bound nucleotide (i.e. GDP-bound (S GDP) and GTP-bound (S GTP)) [1–3]. In the S GTP active state, the G proteins display high affinity for binding downstream effectors, thereby controlling a variety of biological processes such as cell signaling, cell shape, motility and polarity, membrane transport pathways, transcription factor activity as well as vesicular or non-vesicular transport [1–3]. Differently from the small Ras GTPases, heterotrimeric G proteins of the Gα family hold a helical (H) domain in addition to the common GTPase (G) or Ras-like domain. As a Rab (Ras-related in brain) GTPase, Sec4p (hereafter indicated as Sec4) plays regulatory roles in multiple steps of intracellular vesicle trafficking [2]. One Rab can modulate and control an entire stage of transport including vesicle budding, delivery, docking, and fusion [4].

The switch-ON process is under the control of Guanine Nucleotide Exchange Factors (GEFs), which catalyze the exchange of GDP for GTP.

Each G protein family holds its own GEF. The cognate GEF of Sec4 is Sec2p (hereafter indicated as Sec2). GEF action involves a series of fast reaction steps, which lead from a binary GNBP-GDP complex via a trimeric GNBP-GDP-GEF complex to a binary nucleotide-free GNBP-GEF complex, which is stable in the absence of the nucleotide [1,3,5]. This series of reactions is reversed by rebinding of nucleotide, predominantly GTP, because of its higher concentration in the cell [6–8]. In this respect, the GEF acts as a
catalyst to increase the rate at which the equilibrium is pulled from $S^{GDP}$ to $S^{GTP}$. Although Mg$^{2+}$ contributes to the tight binding of nucleotides, GEFs further enhance nucleotide release even in the absence of Mg$^{2+}$ (reviewed in Ref. [1,3,5]). Structure determinations and molecular simulations provided insights into the mechanism of G protein activation. MD simulations on small GTPases from different families revealed a common semi-open conformation similar to the GEF-bound state and suggested a regulatory role of Mg$^{2+}$ [9]. MD simulations on a number of binary and ternary complexes of H-Ras served to infer changes in coordination of Mg$^{2+}$ along the path to G protein activation [10].

The switch-OFF process is regulated by GTPase activating proteins (GAPs), which enhance the relatively slow intrinsic GTPase activity of G proteins. A number of simulation studies investigated nucleotide-dependent conformational states and the reaction path of GTP hydrolysis in small GTPases [15–19].

Like all Ras GTPases, Sec4 holds an αβα three-layer sandwich architecture and a Rossmann fold (Figs. 1 and 2). The helices α1 and α5 lie on the side of the parallel β-sheet, whereas α2, α3, and α4 lie on the opposite side. The nucleotide docks into a binding site contributed by the β1/α1, β1/β2 (α1/β2 in the Gx proteins), β3/α2, β5/α4, and β6/α5 loops (i.e. G boxes 1–5 (G1-G5) Fig. 2 AB). Sequence conservation over the Ras superfamily resides just in the G boxes, also defined as ultra-conserved regions (Fig. 1). G2 and G3 are also defined as switches I and II (SWI and SWII, respectively) and the β2-β3 hairpin preceded and followed by the switches is named inter-switch. Notably, SWII includes also the α2-helix (Fig. 1). Computational experiments suggested that the β1/β4 interface divides the Ras-like domain into two dynamically distinct lobes: lobe 1 (the N-terminal half of the domain) that includes the β1-β3 strands, the P loop, and the two switches; and lobe 2 (the C-terminal half) that includes the β4-β6 strands and the α3-α5 helices [15,21].

The crystal structures of the $S^{GDP}$ and $S^{GTP}$ states of Sec4, respectively bound to Co$^{2+}$ and Mg$^{2+}$ (PDB: 1G16 and 1G17, Fig. 2A-B) [22], show clear differences in the switch regions. Indeed, the turn to the active state induces two-residue lengthening of the N- and C-terminations of β2 and β3, respectively, and a one-turn extension of α2/SWII (Fig. 2A-B). As a consequence, in $S^{GTP}$ the two switches become interacting forming the interaction Site 1 (155/SWI-I57/SWI-6,89/SWI-15) and Site 2 (155/SWI-I57/SWI-5,153/SWI-3,F82/SWI-3) (Fig. 2B). In both $S^{GDP}$ and $S^{GTP}$ crystal structures the nucleotide is caged by the G-boxes: whereas G1, G2, and G3 interact with phosphates and Mg$^{2+}$, G4, G5, and G6 interact with the guanine base. G1 (P-loop or Walker A motif GxxxGKS/T), G2 (or SWII), and G3 (participating in SWII) hold conserved glycines and essentially employ the backbone NH-groups to bind phosphates. The aspartate of DxxG (i.e. DGH1) in G3 (also called the Walker B motif) makes a water-mediated contact with Mg$^{2+}$, which is critically required for tight nucleotide binding and GTP hydrolysis in Ras GTPases. Together with the P-loop that interacts with the β-phosphate, the most important contributions to tight binding result from the interactions of the guanine base with G4 and G5. In this respect, specificity for guanine is realized by the aspartate side chain in position G4:4, which usually forms bifurcated H-bonds with the NH-group at position 1 and the NH$_2$-substituent at C2 of the guanine ring (Fig. 2A-B). The guanine ring makes also σ-π orthogonal interactions with F45 in the loop between α1 and SWI (Fig. 2A-B), whereas the ribose O3’ is engaged in H-bond with the backbone carbonyl oxygen atom of P47 in the same loop.

Nucleotide exchange requires, respectively, loss and gain of such interactions when GDP exits and GTP enters with the aid of the GEF domain of Sec2. In spite of the rather fast rate of nucleotide dissociation from Sec4, Sec2 is still capable to high efficiently stimulate nucleotide release [23].

The N-terminal domain of Sec2 is a prototype for a structurally distinct family of Rab GEFs, which forms a single 220 Å-long parallel coiled-coil (PDB: 2E7S [24], Fig. 2C). Insights into the mechanism of GDP release come from the crystal structures of PO4-bound and APO Sec4 in complex with the GEF domain of Sec2 (PDB: 2EQB [25] and 2OY2 [26], respectively, Fig. 2D,E). The 2EQB complex with the P-loop of Sec4 bound to the phosphate ion represents an intermediary state compared to the 2OY2 GDP-depleted state of the G protein. Furthermore, the crystal structure of Sec2 in complex with the GDP-bound Sec4-S29V mutant, characterized by reduction of both intrinsic and stimulated nucleotide exchange (PDB: 4ZDW [27]), may represent an intermediary state preceding GDE release. Remarkably, the structure of such Sec4 mutant does not hold Mg$^{2+}$ indicating that such cation may be not essential for tight nucleotide binding to Sec4. In the crystal structures of Sec4-Sec2 complexes, Sec4 residues (45–58 and 81–89) in and around SWI and SWII all participate in the hydrophobic interface with the GEF. Such interface is contributed by: F49/SWI-I55/SWI-7,W74/SWI-6, I53/SWI-5, F57/SWI-3, F74/SWI-5, I85/SWI-6, and Y89/SWI-10 of Sec4 and F109, T105, and V101 from what we label as chain A of Sec2 (corresponding to chain B in 2E7S, 2EQB, and 2OY2 structures) as well as L104 and L108 from what we label as chain B of Sec2 (corresponding to chain A in 2E7S and 2OY2 and to chain C in 2EQB, Fig. 2C,D). Long-range salt bridge interactions at the interface include K22/SWI-I103/SWII-D103/SWII-D103/Sec2 and D43/K121/SWII (Fig. 2D). Remarkably, compared to the $S^{GDP}$ state, F45 looses the interaction with the guanine ring of GDP and interacts with both K121/Sec2 and Y124/Sec2 (Fig. 2A, D,E). Sec2 favors the establishment of the interactions between the two Sec4 switches leading to formation of the $S^{GTP}$ Site 1, whereas the amino acids in $S^{GTP}$ Site 2 participate in the interaction with Sec2 (Fig. 2B,D,E). Along the same line, Sec2 also favors lengthening of β2, β3, and α2 in preparation of the $S^{GTP}$ state (Fig. 2B,D,E).

Collectively, structure determinations provided invaluable pictures of the inactive $S^{GDP}$, active $S^{GTP}$, Sec2-bounded $S^{APO}$, and a couple of Sec2-bounded intermediary states of Sec4. However, significant insights into the nucleotide exchange process may derive from molecular dynamics (MD) simulations. By simulating the dynamics of isolated Sec2 and of Sec4 in: a) complex with GDP-Mg$^{2+}$ ($S^{GDP}$), b) complex with GTP-Mg$^{2+}$ ($S^{GTP}$), c) a modeled quaternary complex with Sec2 and GDP-Mg$^{2+}$ ($S^{GDP}$), and d) a binary complex with Sec2, the present study addresses a number of questions concerning the intrinsic and stimulated dynamics of Sec2 and Sec4 as well as the chain of structural deformations leading to GEF-assisted activation of the Rab GTPase.

2. Methods

2.1. MD simulations

2.1.1. The input systems

Sub-microsecond and microsecond MD simulations in explicit water were carried out by the GROMACS (v.4.0.7 [28]) package on the following systems: a) the crystal structures of the $S^{GDP}$ (PDB: 1G16, Sec4 [47]) and $S^{GTP}$ (PDB: 1G17, Sec4 [47]) states of Sec4 [22] with nucleotide, Mg$^{2+}$ ion (in $S^{GDP}$ Co$^{2+}$ was replaced by Mg$^{2+}$), and coordinating water molecules; b) the crystal structure of nucleotide-free Sec4 in complex with Sec2 (following the elimination of the phosphate ion mimicking the β-phosphate of GDP in

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In this respect, the crystallographic structures of Sec4 GDP and unbound Sec2 (PDB: 1G16, chain C; 2E7S, chains A and B, residues 50–142, respectively), i.e. unbound-unbound docking, or Sec4 GDP and Sec2 extracted from the complex with Sec4 APO (PDB: 2OCY), i.e. unbound-bound docking, were used as inputs of protein–protein docking following the default RosettaDock protocol [29,30]. As for the unbound-unbound docking, the two proteins were pre-oriented by fitting each protein on the corresponding components of the Sec4-Sec2 complex (PDB: 2EQB, selected for the fitting because of the most complete structure of Sec4). As for the unbound-bound docking, only Sec4-GDP was pre-oriented upon fitting the structure onto Sec4 from the Sec2-Sec4 complex. In general, as for Sec2 fitting, either all or only the interfacial residues were considered, whereas for Sec4 fitting, all residues except the two switch regions were considered. Before composing the input assemblies, the side-chain conformations of each docking partner were optimized by the “pre-packing” procedure [30]. For each input system, 1000 independent docking runs were done by randomly perturbing the initial orientation of the G protein [31]. For side-chain packing, extra-rotamers were used for all \( \chi_1 \) dihedrals and for the \( \chi_2 \) dihedrals of aromatic residues, including also rotamers from the unbound structures [31,32]. The best 200 poses, ranked according to the interface energy score [33], were retained from each docking assembly and evaluated for the number of interfacial native contacts, defined as those with at least one atom pair within 4 Å in the crystallographic complexes 2EQB or 2OCY. The top poses were subjected to interfacial residue-based clustering by means of Wordom [34], using a Cα-atom RMSD cutoff of 3 Å. Visual inspection of the top-scoring, most representative complex structures was instrumental in selecting 52 input complexes for MD simulations.

### 2.1.2. Simulation setup

Simulations were carried out by the GROMACS 4 (4.0.7) simulation package [28]. The AMBER03 all-atom force field was employed [35], using the TIP3P water model to explicitly describe the solvent. Available topologies and parameters for GDP and GTP for the AMBER force field were used [36]. Depending on the dimension of the system, a variable number of Na\(^+\) and Cl\(^-\) ions were introduced to neutralize the system. Each simulation system was subjected to Periodic Boundary Conditions (PBC) with different unit cell boxes. In this respect, for isolated GTPases, an octahedric box with a distance of 12 Å between solute and boundaries was chosen, whereas for both the crystallographic and predicted Sec4-Sec2 complexes, 10 5 3 (xyz) box was used and aligned to the complex main axis. Such a box approximated the elongated shape of the Sec2 coiled-coil and reduced the number of solvating waters. To keep the principal axis of the complex aligned to the box main axis, translations and rotations of the mass center were removed during each simulation step.

All input structures were subjected to energy minimization using 500 steps of Steepest Descent, with an energy tolerance for convergence estimation of 1000 (kJ mol\(^{-1}\) nm\(^{-1}\)), followed by 500 steps of Conjugate Gradient, with an energy tolerance of 500 (kJ mol\(^{-1}\) nm\(^{-1}\)). The systems were then equilibrated for 4 ns of isothermal-isobaric (\( T = 300 \text{ K}, P = 1 \text{ atm} \)) MD simulations, restraining the position of all protein main-chain and C\( \beta \) atoms, nucleotide,
Mg$^{2+}$ and coordinating water molecules. The v-rescale thermostat [37] was employed to keep the system at a constant temperature of 300 K, using a coupling constant ($\tau_s$) of 0.1 ps. The pressure of the system was kept fixed at 1 atm, using the Berendsen weak coupling algorithm [38] with a coupling constant ($\tau_p$) of 1 ps. The Particle Mesh Ewald (PME) method was employed to compute electrostatic interactions, using a cutoff of 14 Å. Short-range repulsive and attractive interactions were computed using a Lennard-Jones potential with a cutoff of 12 Å. The LINCS algorithm [39] was used to allow for an integration time step of 2 fs by the leap-frog algorithm. The pre-equilibrated systems were then released for 1 ns prior to unrestrained MD simulations in the NPT ensemble (T = 300 K, P = 1 atm). For each crystallographic input structure, 5 independent replicas of 100 ns each were achieved by randomizing initial velocities.

For the S$^{GDP}$ state, 52 Rosetta docking poses were subjected to 100 ns of unrestrained MD simulations (NPT ensemble). The trajectories showing some rearrangement towards the GEF-bound state were extended up to 1 μs. Herein only the analysis of the best conversion is shown. It is worth recalling that in the relative input complex, Sec4 is the crystallographic S$^{GDP}$ state (PDB: 1G16), whereas Sec2 comes from the crystallographic complex with Sec4 APO (PDB: 2OCY). All simulations arising from unbound-unbound input complexes (i.e. with both Sec4 and Sec2 from their unbound inactive crystallographic structures, 1G16 and 2E7S, respectively) did not produce acceptable transitions.

The time series of Cα-Root Mean Square Deviations (Cα-RMSDs) concerning S$^{GDP}$, S$^{GTP}$, and S$^{AP0}$ are shown in Supplementary Fig. S1.
2.2. MD analyses

Comparative analyses of the MD trajectories of the different Sec4 and Sec2 states were carried out by the in-house software Wordom [34] and PSNtools [40].

Incidentally, to save clarity, we renamed as chain A what is called chain B in 2E7S, 2EQB, and 2OCY structures, whereas we renamed as chain B what is called chain A in 2E7S and 2OCY and chain C in 2EQB.

2.2.1. Analysis of flexibility

MD trajectories were subjected to a variety of analyses aimed at inferring the intrinsic flexibility of the systems (i.e. Cx-RMSDs, Cx-Root Mean Square Fluctuations (Cx-RMSFs), force constants [41–43], overall fluctuations [44], collective motions by Principal Component Analysis (PCA) [45], as well as functional mode Analysis (FMA) [46]. As for FMA, it consists in searching for possible correlations between structural features and essential motions, by using the Pearson Coefficient. In this framework, several size/shape and intermolecular interaction descriptors were correlated with linear combinations of a variable number of PCs. All these analyses were performed by means of the Wordom software [34]. All structural descriptors were computed by Wordom as well [34].

For a selection of computational indices, a detailed description is shown as follows.

As for force constants, the mechanical properties at the single residue level were inferred by computing the residue force constants through the analysis of the mean fluctuation of the mean residue level were inferred by computing the residue force constants through the analysis of the mean fluctuation of the mean residue level [34].

\[
\kappa_i = \frac{3k_BT}{\left< (d_i - \left< d_i \right> )^2 \right>}
\]  

(1)

where \(d_i\) is the mean defined distance above, \(\left< \cdot \right>\) denotes the average over the simulation, \(k_B\) is the Boltzmann constant, and \(T\) is the simulation temperature.

Protein dynamics was also investigated by computation of the overall fluctuation index \(\Theta\), which is a measure of the intrinsic flexibility of the whole protein (or of a given sub-set of residues) and is proportional to the extent of conformational space explored in a simulation [44]. \(\Theta\) is defined as root mean distance variance of each atom pair and is calculated by the following equation:

\[
\Theta_{AB} = \sqrt{\frac{\sum_{i=1}^{N} \sum_{j=1}^{M} \frac{\sum_{k=1}^{F} (d_{ij}^k - \bar{d}_{ij}^k)^2}{F}}{N \times M}}
\]  

(2)

where \(A\) and \(B\) are two sets of residues, \(N\) and \(M\) are the total number of atoms in set \(A\) and set \(B\), respectively, and \(F\) is the total number of trajectory frames. Furthermore, \(d_{ij}^k\) is the distance between atom \(i\) from residue set \(A\) and atom \(j\) from residue set \(B\) in the \(k^{th}\) frame while \(\bar{d}_{ij}\) is the average distance between the same two atoms. Here we considered all Cx-atoms in each helix, strand, G box, switches, and the C1 atom of the nucleotide to define the sets of amino acid residues for mean pairwise distance variance computation.

Comparisons of the overall fluctuations of such residue sets between \(\text{SCDP}\) and \(\text{STGP}\) between \(\text{SAPD}\) and \(\text{SCDP}\) or \(\text{SGDP}\), and between \(\text{SCDP}\) and \(\text{SGDP}\) were computed according to the following formulas:

\[
\Delta \Theta\% = 100 - \frac{\Theta_{\text{min}}}{\Theta_{\text{max}}} \times 100
\]  

(3)

where \(\Theta_{\text{min}}\) and \(\Theta_{\text{max}}\) indicate, respectively, the states with minimal and maximal \(\Theta\) values.

As for PCA of the different Sec4 or Sec4-Sec2 states, the covariance matrices were built on the Cx-atoms of the isolated MD trajectories by fitting all Cx-atoms. The same fitting strategy was employed to compute the Cx-RMSF.

2.2.2. Analysis of the structural communication by the protein structure networks

The structural communication was investigated by the protein structure network (PSN) analysis implemented in the PSNtools [40]. PSN analysis is a product of graph theory applied to protein structures [47]. The methodological approach for computing the structure graph and the shortest communication pathways has been described in a number of research articles [40,48,49].

Shortly, in a PSN, each linked residue (e.g. amino acid, nucleotide, small molecules, ion, etc) is a node [50]. Links form if the non-covalent interaction strength between pairs of nodes equals or overcomes a cutoff \((I_{\text{min}})\). Such interaction strength, expressed as a percentage, is computed by the equation (4) below:

\[
I_{ij} = \frac{\text{nij}}{\sqrt{N_i N_j}} \times 100
\]  

(4)

where \(I_{ij}\) is the percentage interaction between residues \(i\) and \(j\); \(n_{ij}\) is the number of heavy atom–atom pairs between the side chains of residues \(i\) and \(j\) within a distance cutoff (4.5 Å); \(N_i\) and \(N_j\) are normalization factors for residue types \(i\) and \(j\), which account for their propensities to make contacts with surrounding residues [47,51].

As for the normalization factors, the PSNtools software employs an internal database holding the normalization factors for the 20 standard amino acids and the 8 standard nucleotides (i.e. dA, dG, dC, dT, A, G, C, and U), as well as for more than 35,000 molecules (e.g. small molecules, lipids, sugars, etc) and ions extracted from all the structures deposited to date in the PDB. The normalization factors for GDP, GTP, and Mg\(^{2+}\) employed in this investigation are 246.47, 248.84, and 19.76, respectively.

Thus, the interaction strengths \((I_{ij})\) are computed for all node pairs. At a given interaction strength cutoff, \(I_{\text{min}}\), any residue pair \(i-j\) for which \(I_{ij} \geq I_{\text{min}}\) (see equation Eqn 4) is considered to be interacting and hence is connected. The \(I_{\text{min}}\) cutoff is automatically computed as described in a recent paper [40]. In more detail, the \(I_{\text{min}}\) employed for the PSN analysis on MD simulations is the average over all the \(I_{\text{min}}\) computed on each trajectory frame.

Only those links and hubs present in at least 33% of the trajectory frames were defined as stable and considered for further analysis.

To avoid excessive network fragmentation, which would impair the search for shortest communication pathway, all node clusters (i.e. ensembles of nodes connected by at least one link) were iteratively connected by the link with the highest sub-cutoff interaction strength.

The allosteric communication was studied by searching for the shortest communication pathways. A pathway describes how signals are transferred between sites and consists of a set of residues in dynamic contact [52,53]. The procedure for computing the shortest communication pathways, which has been previously described and validated [40,49], is based on Dijkstra’s algorithm [54].

A coarse/global picture of the whole structural communication in the considered system is provided by consensus paths or metaphaths made of the most recurrent links in the path pool (i.e. with a recurrence ≥ 20% in this study) [40].

2.3. Analysis of amino acid conservation

Conservation analysis was performed by using the ConSurf web server (https://consurf.tau.ac.il) [55]. In a typical ConSurf applica-
tion, the query protein is first BLASTed against the UNIPROT-90 database. Redundant homologous sequences are then removed using the CD-HIT clustering method. The resulting sequences are next aligned and the generated multiple sequence alignment (MSA) is used to reconstruct a phylogenetic tree. Given the tree and the MSA, the Rate4Site algorithm is used to calculate position-specific evolutionary rates under an empirical Bayesian methodology. The rates are normalized and grouped into nine conservation grades 1-through-9, where 1 includes the most rapidly evolving positions, 5 includes positions of intermediate rates, and 9 includes the most evolutionarily conserved positions [55]. We employed the sequences in the crystal structures of Sec4 (PDB: 1G16) and Sec2 (PDB: 2EQB) to run the ConSurf analysis.

3. Results

3.1. Functional dynamics of Sec2, Sec4, and their mutual association

The main goals of the present investigation are to explore the intrinsic dynamics of: a) the Sec2 GEF homodimer, b) Sec4 in ternary complex with hydrated Mg$^{2+}$ and either GDP (S$^{\text{GD}}$) or GTP (S$^{\text{GT}}$), c) APO-Sec4 in complex with Sec2 starting from the PO$^{4-}$ bound intermediary state of the Rab protein (S$^{\text{APO}}$), and d) the Sec2-induced transition of Sec4 from S$^{\text{GD}}$ to S$^{\text{APO}}$. The goals were accomplished by comparative MD simulations followed by several structural analyses by established approaches already employed to investigate dynamics and structural communication in a large number of Ras GTPases [14,56–61].

3.1.1. The intrinsic dynamics of Sec2 suggests conformation selection by Sec4

Since the minimal Sec2 sequence required for formation of a stable complex with Sec4 and to exert a GEF action is the region 51–142 [27], all simulations considered only the 50–142 Sec2 segment.

Clear structural differences emerge from comparisons of the free (i.e. 2E7S [24]) and Sec2-bound states (i.e. the crystal structures 2EQB, 2OYC, and 4ZDW) of Sec2. The relative orientation of the two helices around the Sec4 binding site (100–121 region) is conserved in all the 10 Sec2$^{11-160}$ dimers in the crystallographically asymmetric unit of Sec2$^{31-160}$ (i.e. 2E7S). Such orientation changes in the complex with Sec4. Indeed, in free Sec2 the two helices cross by 20° in the 100–121 region (inter-main-axis angle = 21.67° for chains A and B of 2E7S, Fig. 3A), whereas in the complexes with Sec4 the two helices are almost parallel (inter-main-axis angle = 7.54° for 2OYC (Fig. 3A) and 9.43° for 2EQB). Other remarkable structural changes between free and Sec4-bound Sec2 concern the patterns of inter-helix interactions in the 100–80 region, which reflects on changes in inter-helical distances.

In detail, in free Sec2 (i.e. Sec2$^{2E7S}$) the two R87 side chains hold almost extended conformations and point towards the main axis of the opposite helix, R87$^{2E7S}$ making H-bond with N84$^{2E7S}$ and R87$^{2OYC}$ making a salt bridge with E91$^{2E7S}$ (Fig. 3A).

Remarkably, in the crystallographic complexes with Sec4 (e.g. Sec2$^{2OYC}$), the two arginines make salt bridges respectively with D83 on the opposite helix, favoring the approaching of the helix main axes (Fig. 3A). Furthermore, in free Sec2, A94$^{2E7S}$ and A99$^{2E7S}$ are distant, being instead interacting in the bound state. Moreover, F94$^{2E7S}$ makes contacts with both L104$^{2E7S}$ and L108$^{2E7S}$, contributing to the coiled-coil interface (Fig. 3A).

Very interestingly, MD simulations of free Sec2, starting from the 2E7S structure (chains A and B), show that the majority of frames hold the two helices almost parallel around the Sec4 binding site (region 100–121, Fig. 3A,B), the median angle between the helix main axes being 7.04 ± 4.24° (Fig. 3B). Along this line, the coiled-coil interface tends to approximate the one of crystallographic Sec4-bound Sec2, the frame closest to Sec2$^{2OYC}$, Sec2$^{MD}$, holding a Cx-atom RMSD 1.82 Å from Sec2$^{2E7S}$ input structure and equal to 0.90 Å from Sec2$^{2OYC}$ (see the left and right insets of Fig. 3B). Remarkably, side-chain interactions at the coiled-coil interface overlap as well between Sec2$^{MD}$ and Sec2$^{2OYC}$ (Fig. 3A). Along the same line, the pattern of inter-helical distances along the 50–142 sequence diverges for Sec2$^{E7S}$ and Sec2$^{2OYC}$ crystal structures, whereas it overlaps extraordinarily well for Sec2$^{MD}$ and Sec2$^{2OYC}$ (Fig. 3C).

Consistently, the distributions of the first five PCA projections clearly show that isolated Sec2 (i.e. Sec2$^{MD}$, Figure S2) explores a wider essential sub-space than Sec2 bound to Sec4 (i.e. Sec2$^{APO}$, Figure S2). In all pairwise combinations of PCs, a partial overlap occurs between the subspaces explored by Sec2$^{MD}$ and Sec2$^{APO}$ (Figure S2).

Collectively, these results unequivocally indicate that free Sec2 is able to adopt a Sec4-bound state even in the absence of Sec4, thus suggesting that the G protein likely selects a pre-formed Sec2 conformation prone to form a productive complex.

3.1.2. Nucleotide and Sec2-GEF affect the intrinsic flexibility of Sec4

The crystal structures of Sec4 provided significant insights into the structural features of the functionally different states of the G protein. The transition from the inactive S$^{\text{GD}}$ (Sec4$^{1G16}$ structure, Fig. 2A) to the active S$^{\text{GT}}$ (Sec4$^{1G17}$ structure, Fig. 2B) involves folding of SWII in a longer α-helix and a re-organization of the interface between the two switches with formation of two interaction sites in S$^{\text{TF}}$ (Fig. 2AB), Such process passes through the interaction of both switches with Sec2, which leads to the GEF-bound S$^{\text{PO}}$ state of the G-protein (Sec4$^{2OYC}$ structure, Fig. 2E). Early stages in nucleotide release are represented by the crystal structures of Sec4 bound to both Sec2 and GDP (Sec4$^{3ZDW}$) or to Sec2 and PO$^{4-}$ (Sec4$^{2EQB}$, Fig. 2D).

To investigate the intrinsic dynamics of S$^{\text{GD}}$, S$^{\text{GT}}$, and S$^{\text{PO}}$ states the crystal structures Sec4$^{1G16}$, Sec4$^{1G17}$, and Sec4$^{2EQB}$ deprived of PO$^{4-}$ were used as inputs of MD simulations. Exploration of the conformational ensembles associated to the three Sec4 states was complemented by simulation of a S$^{\text{GD}}$-transition to a Sec2-bound form (i.e. S$^{\text{GD}}$) by using a modeled complex between Sec4$^{1G16}$ and Sec2$^{2OYC}$ as an input (see the Experimental Section).

Cα-RMSFs remark evidence from computational experiments that in S$^{\text{GD}}$ and S$^{\text{GT}}$ of Ras GTPases flexibility essentially concerns the SWI (49-55 Sec4) and SWII (75-89 Sec4) regions, which are more flexible in S$^{\text{GD}}$ than S$^{\text{GT}}$ (Fig. 4) [56,59,61]. In the predicted S$^{\text{PO}}$, the flexibility of SWII is the highest compared to the other states. Unique feature to S$^{\text{PO}}$ is the higher flexibility of the α1/β1-loop or P-loop (G1), α2, α3, G4, and G5 compared to all other forms (Fig. 4), whereas SWI shows the lowest flexibility likely due to the intermolecular interactions with Sec2. In general, helices show higher flexibility than β-strands due to the α:α-sandwich architecture that buries the β-sheet by the two layers of α-helices. The higher flexibility of the G-boxes G1, G4, and G5 in S$^{\text{PO}}$ is likely due to the lack of the nucleotide and is in line with the observations that loops in the Sec4 nucleotide-binding pocket are disordered as a consequence of nucleotide removal [26].

The mechanical properties of the considered systems at a single-residue level were computed by inferring the force constant from the mean fluctuations of the mean distance from each residue with respect to the rest of the protein structure (see the Experimental Section) [41–43]. Force constants are almost complementary to Cx-RMSFs, in the sense that major rigidity (i.e. increase in force constants) concerns regions with lower flexibility (Fig. 5AD). In this respect, the most rigid regions are, as expected, the β-strands participating in the parallel β-sheet with more prominence.
to β3 and β4, which are adjacent in space and occupy inner positions in the β-sheet. This behavior is due to the three-layer αβα architecture of the Rossmann fold, independently of the functional state. Highly conserved amino acids (i.e. with ConSurf grades ≥ 7) tend to hold force constants above the average, consistent with the demonstrated relationships between amino acid mechanical rigidity, evolutionary conservation, and belonging to the folding nucleus in 70 protein structures from 14 protein families [62]. In this respect, the β4-strand, which marks the beginning of the C-terminal half of the G protein, shown to be dynamically distinct from the N-terminal one [59], reaches the highest rigidity level in all forms, SAPO showing a lower level of rigidity in its C-terminal half, excluding β5, than the other forms (Fig. 5). Indeed, pairwise comparisons between the different states show that, in $S^{\text{GDP}}$, SWI, SWII, inter-switch, and nucleotide display higher overall fluctuations than they do in $S^{\text{GTP}}$ (Fig. 6A). In $S^{\text{GDP}}$, SWI and inter-switch fluctuate in tandem with the other Sec4 portions more than they do in the predicted complex between Sec4 1G16 and Sec2 2OCY (Fig. 6B). Remarkably, $S^{\text{GDP}}$' is closer in overall fluctuations to $S^{\text{APO}}$ than to $S^{\text{GDP}}$ (Fig. 6B-D), not only because both $S^{\text{GDP}}$ and $S^{\text{APO}}$ concern GEF-bound forms, but also because $S^{\text{GDP}}$ likely represent an evolution of $S^{\text{GDP}}$ towards the nucleotide-depleted form. In both the simulated GEF-bound forms, the whole Sec2 shows overall fluctuations with all Sec4 portions, which are higher in $S^{\text{GDP}}$ than $S^{\text{APO}}$, suggesting that the former complex is more dynamic and less stable than the latter (Fig. 5D).

Collectively, despite the presence of bound GDP and Mg$^{2+}$, the flexibility of Sec4 in $S^{\text{GDP}}$ resembles that of $S^{\text{APO}}$ more than $S^{\text{GDP}}$, indicating that $S^{\text{GDP}}$ expresses an evolution towards nucleotide release. The nucleotide-depleted $S^{\text{APO}}$ form shows enhanced flexibility of the P-loop and the C-terminal half than the other forms, which may be linked to the structural rearrangements preventing GDP re-binding [25].

Fig. 3. Intrinsic dynamics of Sec2. A, top. The helix main-axis orientations of the 100–121 region of Sec2 crystallized in its isolated state (Sec2 2E7S), in complex with Sec4 (Sec2 2OCY), or simulated in the isolated state (Sec2 MD) are shown. Helices A and B are shown as yellow and aquamarine cartoons, respectively. As for Sec2 MD, the frame closest in Ca-RMSD to Sec2 2OCY is shown. A, bottom. A different view of sections 81–110 of the two helices is shown for Sec2 2E7S, Sec2 2OCY, and Sec2 MD. B. The superimposed structures of Sec2 2OCY (violet) with Sec2 2E7S (left orange) and with Sec2 MD (right orange) are shown. The orange boxplot reports the median value and the standard deviation of the angle between the helix-main axes in the region 100–121. The two black lines extending from the box indicate the variability outside the upper and lower quartiles and the two black dots represent the largest and lowest values of the angle. C. The distances between the a-helix axes of Sec2, as a function of residue number, is calculated as the distance between the projections of the geometric backbone-atom center of each residue on the local helix main axis. For each residue, the local main axis of each helix is recalculated using the backbone coordinates of the residue and of the six preceding and following residues in the primary sequence. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
The analysis of Sec4 crystal structures allowed us to define a number of structural hallmarks of the different functional states. Such hallmarks, computed on the trajectory frames, contributed to infer hypotheses on the mechanism of nucleotide exchange.

In this respect, we tested a huge number of descriptors finally selecting the following: a) the distance between the Cα-atom of G32 and the centroid of the F49 side chain; b) the angle among the Cα-atoms of R39, S34, and K44; and d) the angle between the main axes of the α2 and α3 helices (by selecting, respectively, the amino acid ranges 86–91 and 113–118 for S GDP, S GDP, and S GDP′, and 82–91 and 112–122 for S APO) (Fig. 7).

Whereas the G32-F49 distance and the angle between the three Cα-atoms describe the deformations of P-loop (G1) and SWI (G2), the angle between α2 and α3 describes deformations of SWII (G3) associated with changes in G protein functional state. Indeed, all those descriptors increase ongoing from the S GDP or S GTP isolated states of Sec4 to its Sec2-bound forms. In particular, in Sec2-bound forms the main axes of α2 and α3 become almost orthogonal following rotation of α2, while participating in the interface with Sec2. The pairwise combinations of the three indices computed on the trajectory frames were effective in distinguishing the different states of Sec4 and highlight the tendency of the conformational ensembles to deviate from the input structure (Fig. 8A-D). In detail, independently of the index, the population-surfaces show the basins of the Sec2-bound forms of Sec4 (S GDP and S GTP) clearly separated from the unbound ones and shifted towards higher values of all three indices (Fig. 8). S GDP and S GTP show more than one basin deviating from the position of the crystallographic input structure. This behavior that is emphasized in S GDP compared to S GTP, is due to the intrinsic flexibility the two switches. Remarkably, the most
relevant deviation from the input structure concerns $S^{\text{GDP}}$ since the Sec4 input structure, i.e. the crystallographic structure Sec4$^{1G16}$ (Fig. 8A), during MD simulations evolves towards the crystallographic structures Sec4$^{2EQB}$ or Sec4$^{2OCY}$ (Fig. 8A) as well as towards the conformational ensemble of $S^{\text{APO}}$ (Fig. 8B-D). The single basin of the $S^{\text{APO}}$ conformational ensemble tends to remain close to the input structure (Fig. 8A-D).

The geometrical descriptors listed above worked well in functional mode analysis (FMA) computed on the different states of Sec4. Indeed, all of them correlated with the collective atomic motions describing the essential subspace (ES) of Sec4 in different functional states. In general, ES is contributed by a variable number of eigenvectors whose associated eigenvalues account for 90% of the total variance of the Cα-displacements in a trajectory.

In this study for all considered states of Sec4, the R39-S34-K44 and α2/α3 angles gave correlations $R^2 \geq 0.93$ (validation set) and $R^2 \geq 0.97$ (model set) with a combination of the first 500 eigenvectors describing ~100% of the total variance of the Cα-displacements (i.e. the ES).

As for the α2/α3 angle, correlations concerning both validation and model sets were relatively low ($R^2 \approx 0.70$) only for $S^{\text{APO}}$. The G32-F49 distance performed slightly worse, since the correlation coefficients $R^2 \geq 0.85$ (validation set) or $R^2 \geq 0.97$ (model set) were achieved for all systems but $S^{\text{GDP}}$, which did not provide correlations above 0.7 $R^2$. For each system, the projections on the Cα-atoms of the weighted combination of the first 500 eigenvectors described the same collective motions. Weighing of eigenvectors was based on their contribution to the FMA correlative model. Therefore, here we show only the projections concerning the collective motions correlated with variations of the R39-S34-K44 angle as exemplar of the collective motions correlated with the other two descriptors (Fig. 9). In general, the weighted combination of the first 500 eigenvectors describes collective motions of SWI. Whereas for $S^{\text{GDP}}$ and $S^{\text{GTP}}$ such modes involve exclusively SWI (Fig. 9A,B), for $S^{\text{GDP}}$ they involve an increasing number of portions, as SWI, SWII, inter-switch, G5, and α5 (Fig. 9D). Remarkably, largest amplitude motions concern SWI, which displaces towards the Sec2 GEF (Fig. 9C). For the $S^{\text{APO}}$ collective motions, even if with

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Fig. 6. Comparisons of the overall flexibility of the different Sec4 states. Pairwise % differences in overall fluctuations between the different portions are expressed as colors in the matrices (see the color legend on the right). The four panels show comparisons between: (A) $S^{\text{GDP}}$ and $S^{\text{GTP}}$, (B) $S^{\text{GDP}}$ and $S^{\text{GDP'}}$, (C) $S^{\text{GDP}}$ and $S^{\text{APO}}$, and (D) $S^{\text{GDP'}}$ and $S^{\text{APO}}$. The sign of the Δ% is relative to the second state listed on each matrix.
lower amplitude compared to S\textsuperscript{GDP}, involve all Sec4 portions, likely linked to the presence of Sec2 and to removal of the PO\textsubscript{4}\textsuperscript{3-} ion mimicking the nucleotide β-phosphate in the Sec4\textsuperscript{2EQB} structure employed as a simulation input. The essential motions in S\textsuperscript{APO} may be linked to the structural rearrangements preventing GDP rebinding [25].

Collectively, MD simulations of the S\textsuperscript{GDP} and S\textsuperscript{GDP'} forms of Sec4 display a strong tendency of SWI to detach from GDP. Such tendency is emphasized by the presence of the Sec2 GEF in the S\textsuperscript{GDP'} trajectory.

3.1.3. Structure and dynamics of Sec4-Sec2 recognition

The analyses of flexibility and FMA suggest that the S\textsuperscript{GDP} form represents an evolution of the inactive G protein towards the nucleotide-depleted form.

To better characterize such form, we performed a structure network analysis of the Sec4-Sec2 interface in the MD trajectories of S\textsuperscript{GDP} and S\textsuperscript{APO}, using the crystal structure of 2OCY as a reference, which represents the Sec2-bound nucleotide-depleted state of Sec4.

In this framework, in the comparative analysis of the Sec4-Sec2 interface, we considered as rather equivalent all Sec4-Sec2 contacts involving at least one amino acid residue at a distance in the range ± 1 - ± 4 residues in primary sequence from the reference residue (i.e. in the 2OCY crystal structure, Table 1, Fig. 10A).

Remarkably, the totality of identical (native) and equivalent nodes at the Sec4-Sec2 interface of S\textsuperscript{GDP'} is 89%, indicating that, during MD simulations the input S\textsuperscript{GDP} Sec4\textsuperscript{1G16} structure, pre-oriented to Sec2 2OCY, evolves towards a Sec2-bound form by conformational changes of the two switches (Fig. 10A). The simulated S\textsuperscript{APO} state upon removal of PO\textsubscript{4}\textsuperscript{3-} from Sec4\textsuperscript{2EQB} shares 79% of the native interface links with the 2OCY crystal structure (Fig. 10B, Table 1). This result is in line with inferences from structure determinations showing that the evolution from the PO\textsubscript{4}\textsuperscript{3-}–bound 2EQB intermediate to the APO 2OCY complex involves conformational rearrangements of P-loop and other Sec4 regions, while keeping invariant the Sec4-Sec2 interface, which involves the two switch regions of the G protein [25].

Similarities in structure communication were inferred also from the metapaths computed on the S\textsuperscript{GDP} and S\textsuperscript{APO} trajectories.

Fig. 7. Structural hallmarks of Sec4 states. The distance between the side-chain centroid of G32\textsuperscript{G1:6} and F49\textsuperscript{G2:1} (purple), the angle among the C\textalpha{}-atoms of R39\textsuperscript{A1:7}, S34\textsuperscript{G1:8}, and K44 (orange), and the angle between the main axes of the \textalpha{}2 and \textalpha{}3 helices (yellow) are shown on the structures of (A) S\textsuperscript{GDP} (green), (B) S\textsuperscript{GTP} (red), (C) S\textsuperscript{GDP'} (marine), and (D) S\textsuperscript{APO} (grey) closest in C\textalpha{}-RMSD to the average structure from the relative MD trajectory. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
Fig. 8. Distributions of selected structural hallmarks of Sec4 states. A. The three geometrical indices: (a) the distance between the side-chain centroid of G2G16 and F49G21, (b) the angle among the Cα-atoms of R39G17, S34G18, and K44, and (c) the angle between the main axes of the α2 and α3 helices are shown on the crystal structures of Sec4G16, Sec4G17, Sec4G28, and Sec4G20. B-D. The distribution surfaces of the conformational ensembles according to pairs of the above defined descriptors are shown for SGDP (green), SGTP (red), SGDP' (marine), and SAPO (grey). Dots refer to the structures used as inputs of MD simulations. For each surface plot, the representative structures of the main basins for each Sec4 simulated form are shown as cartoons with the pairs of geometrical descriptors drawn. Plots were generated by calculating the multivariate kernel density estimation of the distributions of each pair of descriptors by means of the MATLAB ver. R2018b numerical analysis package with all default parameters. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
Indeed, both metapaths express a communication between the hydrophobic interface, which involves highly conserved amino acids (i.e. with ConSurf grades ≥ 7), and the N-terminal regions of Sec2, passing through a chain of alternate anionic and cationic amino acids with i, i + 4 or i, i + 3 patterns on Sec2β (i.e. E93, K89, E85, K82, and D79) (Fig. 10 C,D, Supplementary Table 1 [Table S1]). This suggests that the hydrophobic interface involving SWI and SWII of Sec4 is structurally/dynamically linked to the N-terminal half of Sec2.

The structural coupling between Sec4-Sec2 interface and N-terminal regions of Sec2 reflects on the essential motions inferred from the S GDP' and in S APO trajectories (Fig. 11). The PCA, done upon fitting all Cα-atoms in the complex, indeed shows a strong overlap between the first eigenvectors (PC1) from the two systems (overlap = 0.91). Such vectors essentially describe the bending of the N-terminal half of Sec2 towards Sec4. Whereas in S APO Sec4 contributes poorly to PC1, in S GDP' all residues participate in PC1 indicative of higher mobility of Sec4 in S GDP' than S APO because of the establishment of the inter-protein interface, upon starting from two poorly interacting proteins. Remarkably, in the S APO MD trajectory started by an already formed Sec4-Sec2 interface, which does not contribute to the essential motions of the complex that instead involve the N-terminal half of Sec2 and the inner regions of Sec4 such as the P-loop and the C-terminal half of the G protein. This is in line with the evidences from structure determinations [25] and from the present MD simulations that the evolution from the PO43--bound 2EQB intermediate to the APO 2OCY complex involves deformations in the majority of Sec4 regions but those participating in the interface with Sec2 [25,26]. Some overlap also concerns PC2 of S APO and PC3 of S GDP' (overlap = 0.78, Fig. 11).

It is worth recalling that the length of porcupines describes the maximal atomic displacement, which, however, occurs rarely in the simulation time. Collectively, PSN and essential dynamics analysis, suggest that Sec2 assists both Sec4 deformations leading to nucleotide release and preventing GDP binding. The results also suggest that the S GDP' trajectory describes an evolution from the starting S GDP towards S APO states of Sec4.
3.2. Hypotheses on the early stages of Sec4 activation by Sec2

By simulating the dynamics of isolated Sec2bound and of Sec4 in four different forms the present study addressed a number of questions concerning the intrinsic and stimulated dynamics of Sec2 and Sec4 as well as the chain of structural deformations leading to GEF-assisted activation of the Rab GTPase.

First of all, MD simulations of isolated Sec2bound show that the Sec4-bound form of Sec2 (e.g. Sec220OCY) pre-exists to complex formation (Fig. 3), thus suggesting that Sec4 selects the Sec2 conformation suitable to productive interaction. This finding is also in line with the observation that all unbound-unbound docking simulations between Sec41G16 and Sec22OCY interface link and the interface link in the MD trajectories of S GDP and S APO. Here the numbers from 0 to 6 (i.e. n) mean that in S GDP the link is contributed by identical nodes (i.e. n = 0) or by nodes found at a primary-sequence distance ± n.

| Node1a | Node2a | Pos1b | Cons1c | Cons2d | S0pos | S0gdp |
|--------|--------|-------|--------|--------|-------|-------|
| A18    | A98    | SWII  | 7      | 6      | 0     | 2     |
| A18    | A101   | SWII  | 7      | 8      | 3     | 1     |
| F45    | A121   | 1x1_SWI | 8     | 6      | 0     | 2     |
| F45    | A124   | 1x1_SWI | 8     | 7      | 0     | 1     |
| F49    | B115   | SWII  | 5      | 9      | 0     | 4     |
| F49    | B112   | SWII  | 5      | 9      | 2     | 4     |
| F49    | B111   | SWII  | 5      | 9      | 2     | 4     |
| F49    | B108   | SWII  | 5      | 9      | 0     | 4     |
| F49    | A113   | SWII  | 5      | 9      | 2     | 4     |
| F57    | A106   | SWII  | 8      | 8      | 0     | 3     |
| F57    | A109   | SWII  | 8      | 9      | 0     | 0     |
| F52    | B103   | SWII  | 8      | 7      | 0     | 1     |
| F52    | B100   | SWII  | 8      | 9      | 0     | 1     |
| F52    | B104   | SWII  | 8      | 9      | 0     | 1     |
| F52    | B108   | SWII  | 8      | 9      | 0     | 1     |
| F52    | B100   | SWII  | 7      | 9      | 0     | 2     |
| K22    | A102   | β1    | 9      | 9      | 0     | 0     |
| P47    | B115   | 1x1_SWI | 1     | 9      | 1     | 5     |
| P47    | A113   | 1x1_SWI | 1     | 9      | 0     | 6     |
| P47    | A116   | 1x1_SWI | 1     | 9      | 0     | 4     |
| R81    | B103   | SWII  | 8      | 7      | 0     | 2     |
| R81    | B107   | SWII  | 8      | 9      | 0     | 4     |
| T52    | B111   | SWII  | 9      | 9      | 0     | 1     |
| T52    | B100   | SWII  | 8      | 9      | 0     | 3     |
| T84    | B100   | SWII  | 8      | 9      | 0     | 3     |
| T84    | B106   | SWII  | 8      | 9      | 0     | 3     |
| W74    | A109   | β3    | 9      | 9      | 0     | 6     |
| W74    | A105   | β3    | 9      | 9      | 0     | 4     |
| Y89    | A108   | SWII  | 8      | 9      | 0     | 1     |

aAmino acid residues from Sec4 (Node1) and Sec2 (Node 2, chains A and B) participating in stable links at the interface between Sec4 and Sec2 in the crystallographic complex encoded as 2OCY. bNode position in Sec4. cConservation grades (according to ConSurf) of the linked nodes in Sec4 and Sec2, respectively. dDegree of equivalence between each nodes (i.e. n = 0) or by nodes found at a primary-sequence distance ± n.

In this context, information from structure determination and from MD simulations is exploited to infer a possible temporal hierarchy of Sec2-assisted deformations in the nucleotide cage ultimately leading to GDP release.

In both Sec41G16 and Sec41G17 crystal structures the nucleotide is caged by the G-boxes and the 1x1/SWI loop. While G1, G2, and G3 interact with phosphates and Mg2+, 1x1/SWI loop, G4, and G5 interact with the guanine base. As for the 1x1/SWI loop, F45 makes σ-π orthogonal interactions with the guanine ring, whereas the backbone carbonyl oxygen atom of P47 makes H-bonding interaction with the ribose O3’. Interactions between the P-loop and the β-phosphate and between the guanine ring and both G4 (e.g. the H-bond between the guanine and D136G4:4) and G5 provide the most important contributions to tight binding. The crystal structure of the S29V mutant bound to GDP and Sec2 (PDB: 4ZDW), representing an intermediary state in the path to GDP release, shows still tight interaction between nucleotide and almost all G boxes, while the interactions between nucleotide and 1x1/SWI loop, SWI, and D134G4:2 side-chain are lost (Figs. 12 and 13A-D).

Indeed, the distances between the centroids of the phenyl ring of F45 and of the imidazole ring of guanine (i.e. F45-guanine distance) and between the carbonyl oxygen atom of P47 and the ribose O3’-atom (i.e. P47-ribose distance) are 5.0 Å and 3.3 Å in Sec41G16 (starting structure of the S GDP trajectory) and 4.9 Å and 3.0 Å in Sec41G17 (starting structure of the S GTP trajectory).

In the presence of the Sec2-GEF, e.g. in the 4ZDW Sec4S29V-GDP-Sec2 complex, those distances increase significantly, e.g. to 17.0 Å and 16.0 Å, respectively, as a consequence of the huge detachment of 1x1/SWI loop and SWI from GDP (Fig. 12).

We recall that formation of the S GTP state is also associated with formation of two sites of interactions (Site 1 and Site 2) between the two switches, which are absent in S GDP (Fig. 2).

To infer a tentative temporal hierarchy in nucleotide-cage deformations, we chose the combination of F45-guanine distance and the plane-angle among the Cα-atoms of R39G1, S34G1, and K44, accounting for the deformations of P-loop (G1) and SWI (G2), as descriptors of the nucleotide exchange process (Fig. 12). In this context, the P47-ribose distance works similarly to the F45-guanine distance. Both distance and angle increase in the presence of Sec2 and separate the Sec2-bound from the Sec2-free states of Sec4 (Fig. 12A). Remarkably, in S GDP, the interaction between F45 and the nucleotide tend to be lost even in the absence of Sec2 (Fig. 12A,B and Fig. 13A). Indeed, in the representative structure of one of the two almost equi-populated basins, the F45-guanine distance is 7.7 Å (Fig. 12A), which is higher than the value in the starting crystal structure (Sec41G16, 5.0 Å). In the representative
structure of the main basin concerning the S GDP’ trajectory, the F45-guanine distance increases to 13.6 Å, reaching a maximum of 25.5 Å within the conformational ensemble (Fig. 12 B).

All together, these results show that early deformations in the nucleotide cage concern the α1/SWI loop and SWI. In the structure from MD simulations of S GDP’ closest to the crystallographic Sec4–Sec2 complex (Fig. 13 B), the α1/SWI loop approaches Sec2A driven by the establishment of a persistent D43 Sec4-K121 Sec2 interaction (Fig. 13 B). The residues K44 and F45, in such loop, approach Y124 Sec2A. Another remarkable feature of S GDP’, which is shared with all crystallographic Sec4–Sec2 complexes, is formation of specific interactions between the two switches characterizing Site 1 (I55 SWI:7–W74 b 3:6–Y89 SWII:15, Figs. 2 and 13 B–D). Site 1 and α2, which is part of SWII, participate in the interface with Sec2.

Remarkably, while establishing contacts with Sec2, α2 becomes almost orthogonal to α3. This is another deformation of Sec4 shared by the Sec2-bound forms, which emerges already from the S GDP’ conformational ensemble (Fig. 13 B). However, due to the less structured state of α2 in S GDP’ compared to the Sec4–Sec2 crystallographic complexes, the side chain of Y89 SWII:10 is still directed towards the parallel β-sheet of Sec4 instead of turning and participating in the hydrophobic interface with Sec2 (Figs. 2 and 13 B–D).

Collectively, the amino acids forming a hydrophobic cluster at the Sec4–Sec2 interface are the same in S GDP’ compared to the Sec4–Sec2 crystallographic complexes, the stereochemistry of the interface and the details of the inter-residue interactions are different (Table 1 and Fig. 13 B–D). In S GDP’, the nucleotide completely looses all interactions with α1/SWI loop while retaining all other interactions characterizing the S GDP state (Fig. 13 A, B).

By integrating the information from structure determination and molecular simulations the following mechanistic picture can be drown: 1) Sec4 selects a Sec2 conformation suitable to form the proper interface with Sec2 (Fig. 13); 2) the establishment of the K22 b 1:3–E102 Sec2A salt bridge and hydrophobic interactions between the C-terminal end of SWI and F109 Sec2A, L104 Sec2B, and L108 Sec2B contributes to early anchoring of Sec4 to Sec2 (Fig. 13 A); 3) the α1/SWI loop and SWI, pulled by Sec2 on itself, lose their contacts with GDP (Fig. 13 B–D); 4) Site 1 interactions between SWI and SWII form (Fig. 13 B–D); 5) the main axis of α2 (belonging to SWII) becomes orthogonal to α3 (Fig. 13 B–D); 6) SWII folds in a 3-turn α-helix allowing for completion of the Sec4–Sec2 interface that is largely hydrophobic (Fig. 13 C); 7) the H-bonding interactions between guanine ring and D136 G4:4 breaks possibly destabilizing also the interactions with both G4 and G5 (Fig. 13 C); and 8) final conformational changes of P-loop, SWI and SWII (i.e. G1, G2, and G3) occur leading to GDP release (Fig. 13 D).

The spontaneous acquisition of a Sec4-bound state by Sec2 and all Sec2-assisted deformations of Sec4 listed at points 3–6 above are likely to occur on fast time-scales as they could be observed...
in microsecond MD simulations. In contrast, Sec2-assisted deformations of Sec4 listed at points 7–9 above are likely to require longer time-scales.

4. Discussion

We have been investigating since long functional dynamics and structural communication in Ras GTPases [14,56–60]. PCA and Elastic Network Model-Normal Mode Analysis (ENM-NMA) were exploited to identify the important structural flexibilities that enable proteins in the Ras superfamily to switch between their active and inactive states [21]. The study led to a hypothesis on the evolutionary adaptation of structural deformations by the individual members of the superfamily to fulfill their specialized function. The deformation modes, which allow the Ras GTPases to accomplish their switching function, are conserved along evolution and reside in the N-terminal lobe-1 portions comprising the two switches [21]. These modes lead to functional specialization when associated with evolution-driven deformations of protein portions essentially located in lobe 2, distal from the nucleotide. Consistently, high pressure crystallography indicated that the complex interplay between N-terminal and C-terminal lobes (i.e. effector and allosteric lobes, respectively) is directly linked to Ras cycling [63].

A combination of ENM-NMA with PSN analysis, in a so-called mixed PSN-ENM approach to infer structural communication pathways from single crystallographic structures, served to dissect functional dynamics and structural communication in GEFs specialized for Rho GTPases (RhoGEFs), which hold the DH and PH domains [64]. The study inferred major evolutionary-driven deformations related to the mechanisms adopted by the GEF to prevent Rho binding (i.e. functional specialization linked to auto-inhibition). MD simulations on isolated and RhoGEF-bound RhoA strengthened the scaffolding action of the DH domain, which primarily turns the SWI of RhoA on itself and enhances the flexibility of the nucleotide binding regions, weakening the connections between GDP and G-boxes [61].

In line with computational investigation of RhoGEF action, the present study employed microsecond MD simulation and established analysis tools to infer hypotheses on the mechanism of GDP release from another Ras GTPase, Sec4, belonging to the Rab family. In general, whereas Ras GTPases are structurally very conserved, GEFs are specific for each family and completely structurally unrelated. Exemplar of this is the fact that RhoGEFs belong to the Dbl family and hold the DH and PH domains, whereas Sec2 is made of a coiled-coil helix dimer. Yet, the RhoA and Rab regions deputed to GEF binding and nucleotide release are the same (i.e. the two switch regions and the inter-switch).

Remarkably, we found that Sec2 has an intrinsic ability to adopt the conformation found in the crystallographic complexes with Sec4, thus suggesting that the latter selects and shifts the conformational equilibrium of the GEF towards a pre-existing bound-like conformation rather than inducing it. In contrast, simulations of Sec4 in the S\(_{\text{GDP}}\) state could not find any Sec2-bound conformation, which may be linked, at least in part, to the more complex energy landscape of the Rab protein compared to the cognate GEF. Deformations of the GDP-bound form towards the APO form of Sec4 could be partially observed only in the presence of Sec2. Indeed, MD simulations of early deformations in the transition from S\(_{\text{GDP}}\) to S\(_{\text{APO}}\) led to the S\(_{\text{GDP'}}\) conformational ensemble, characterized by pulling of SWI on Sec2 and consequent displacement of the a1/SWI loop and the whole SWI away from the nucleotide cage. Formation of Sec2-Sec4 interface was accompanied by extensions of both \(\beta_2\) and \(\beta_3\) strands and formation of Site1-interactions between the two switches in preparation of the active state. Preliminary folding and re-orientation towards Sec2 of SWI was also observed. Although productive deformations leading to GDP release require the presence of Sec2, some of those deformations could be already observed in the conformational ensemble characterizing

\[ \text{Fig. 11. Essential motions of the Sec4-Sec2 complexes. The porcupine representations of PC1 and PC3 concerning S}_{\text{GDP'}}\text{ and those of PC1 and PC2 concerning S}_{\text{APO}}\text{ are shown. The two PC1 have a 0.91 overlap, whereas PC3 and PC2 have a 0.78 overlap.} \]
the inactive $S^{\text{GDP}}$ state. Some of them include loss of the F45-guanine interaction that characterizes the crystal structures of isolated Sec4 and formation of Site 1 interactions observed in all crystallographic complexes between Sec4 and Sec2, thus highlighting an intrinsic ability of the Rab protein to lose interactions between nucleotide and $\alpha_1$/SWI loop and SWI. All together these features may be linked, at least in part, to the rather high rate of nucleotide dissociation by Sec4.

Whereas pulling of $\alpha_1$/SWI loop and SWI on Sec2 and formation of Site 1 occur early in the process of GDP release, the establishment of the native hydrophobic Sec2-Sec4 interface participated by SWI and SWII is slower as it requires a reorganization of the interface between SWII and both $\beta_1$ and $\beta_3$, accompanied by extension and re-orientation of $\alpha_2$ (which is part of SWII). Thus, the rearrangement of the hydrophobic interface leading to the final deformations of the G boxes, in particular G1-3, could not be observed in $S^{\text{GDP}}$ trajectory. In the Sec4-Sec2 system, such event is likely linked to the establishment of a structural communication between the hydrophobic interface and the N-terminal portion of the GEF. In this respect, differently from the RhoGEF DH, which essentially acts as a scaffold for RhoA binding, Sec2 seems to dynamically aid Sec4 transformation.

In summary, the results of computational experiments, integrated with structure determinations, suggest that Sec4 selects the suitable conformation of Sec2, by establishing early

Fig. 12. Distributions of structural hallmarks of nucleotide-cage deformation. A. The geometrical indices: R39$^{\alpha_1}$, S34G1$^{\alpha_1}$, and K44, and the distance between F45 centroid and guanine pentatomic-ring centroid (i.e. F45-Nuc distance) are used as coordinates in population surface plots concerning the Sec4 forms $S^{\text{GDP}}$ (green), $S^{\text{GTP}}$ (red), and $S^{\text{GDP}}$ (marine). Dots refer to the structures used as inputs of MD simulations. The magenta dot refers to the crystal structure of GDP-bound S29V mutant of Sec4 in complex with Sec2 (PDB: 4ZDW). On the right side, the representative structures of the main basins for each simulated form of Sec4 are shown as cartoons with the pairs of geometrical descriptors drawn. Cartoons of the 4ZDW crystal structure together with the geometrical descriptors are shown as well. B. Box-plots of the F45-Nuc distance are shown. On the right side, the structures corresponding to the frames with the maximal F45-Nuc distance are shown as cartoons with the pairs of geometrical descriptors drawn. Surface plots were generated by calculating the multivariate kernel density estimation of the distributions of each pair of descriptors by means of the MATLAB ver. R2018b numerical analysis package with all default parameters. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
interactions such as the K22b13-E102Sec2A salt bridge and a provisional hydrophobic interface. This favors Sec2-assisted pulling on itself of the a1/SWI loop and SWI, which loose any contact with GDP. Pulling of a1/SWI loop and SWI, formation of SWI-SWI Site 1 interaction, and rotation of a2 likely occur earlier. Formation of the final hydrophobic interface, which would require final deformations of SWI and SWII, accomplishes later. Disruption of the nucleotide cage would cause firstly the loss of interactions with the guanine ring and secondly the loss of interactions with the phosphates.

In line with the results of high pressure crystallography determinations on Ras, revealing the multiplicity of sub-states with many transient intermediates [63], the dynamics of Sec4 is more complex than that of its GEF, which may relate to the latter acting
as a catalyst. Indeed, the ease in sampling the energy landscape and adopting a bound-like conformation likely favors the catalyzing ability of a Ras GTPase GEF.

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Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.csbj.2022.09.016.

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