Evolutionary conservation of the WASH complex, an actin polymerization machine involved in endosomal fission

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**W**ASH is the Arp2/3 activating protein that is localized at the surface of endosomes, where it induces the formation of branched actin networks. This activity of WASH favors, in collaboration with dynamin, the fission of transport intermediates from endosomes, and hence regulates endosomal trafficking of several cargos. We have purified a novel, stable multiprotein complex containing WASH, the WASH complex, and we examine here the evolutionary conservation of its seven subunits across diverse eukaryotic phyla. This analysis supports the idea that the invention of the WASH complex has involved the incorporation of an independent complex, the CapZ α/β heterodimer, forming the so-called Capping Protein (CP), as illustrated by the yeasts *S. cerevisiae* and *S. pombe*, which possess the CP heterodimer but no other subunits of the WASH complex. The alignments of the orthologous genes that we have generated give a view on the conservation of the different subunits and on their organization into domains. Moreover, we propose here a unique nomenclature for the different subunits to prevent future confusions in the field.

The WASH complex has been purified from human cells by tandem affinity purification.1 WASH was found associated with VPEF, KIAA0196, CP, CapZ, Cdc53. The native WASH protein is found in a large complex characterized by a Stokes’ radius of 88 Å and a sedimentation coefficient of 12.5 S.1 Moreover, depletion of any of the above-mentioned proteins associated with WASH destabilizes WASH. Together these properties are compatible with a stable complex containing one molecule of each of these proteins. One expects to find all the genes encoding the 7 subunits of this molecular machine in species where this organization into a multiprotein complex is conserved. So using the BLAST algorithm, we looked into public databases for proteins homologous to the ones we had purified. Genes encoding subunits of the WASH complex were detected in genomes of animal, fungi and amoeba species. But none of the subunits were detected in bacterial and plant genomes. The orthologous subunits were retrieved and compared to the mouse subunits chosen as a reference instead of the human ones, to avoid the complexity of the human WASH family.2,3

The two subunits forming the CP cap actin filaments and hence blocks filament elongation. The 2 CP subunits, but no other subunits of the WASH complex, were detected in the *S. cerevisiae* and *S. pombe* yeasts. The same was true for other sequenced fungi. Since the amoeba *D. discoideum* possesses most WASH complex subunits, the ancestor of fungi and amoeba likely possessed these genes, suggesting that the WASH complex has been subsequently lost in fungi. In fungi genomes, the occurrence of CP without the WASH complex is in line with a free heterodimer representing the major pool of CP independently of the WASH complex.1,4 The CP heterodimer is also an integral component of the Dynactin complex.4 The many interactions of CP with actin, the WASH complex and the dynactin complex must exert a strong selective pressure on the CP genes, probably explaining

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**Key words:** Arp2/3 complex, endosome, VPEF, KIAA0196, KIAA0196, CP, CapZ, Cdc53

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VPEF, which is anterior and descriptive. The major pool of the WASH complex and hence of VPEF is associated with endosomes. Nonetheless, the WASH complex is recruited to the plasma membrane upon infection with pathogens. WASH is detected at attachment sites of Salmonella, and is involved in the entry of this pathogenic bacterium into the cell. Similarly, VPEF downregulation impairs the entry of Vaccinia virus. A surprising result, however, was that entry of Vaccinia is also blocked by extracellular antibodies directed against VPEF or by soluble VPEF in the culture medium, as if VPEF played the role of an extracellular receptor for the virus. These observations remain to be explained, because the WASH complex is cytosolic and no transmembrane complex and to the N-terminal domain of N-WASP, which complexes N-WASP with WIP family proteins. The rest of WASH, including the VCA domain, is predicted to be unstructured. The VPEF subunit is of major interest, but confusing. VPEF was first characterized as a factor required for Vaccinia virus entry into mammalian cells, hence its name Vaccinia Penetration Factor. Unfortunately, this publication has been overlooked in the two recent publications reporting VPEF association with WASH. We used the name of the cDNA, KIAA0592, to refer to this protein. Gomez and Billadeau used the name FAM21, standing for FAMilly number 21 coined in automatic annotations of protein families. We recommend to use the name VPEF, which is anterior and descriptive. The major pool of the WASH complex and hence of VPEF is associated with endosomes. Nonetheless, the WASH complex is recruited to the plasma membrane upon infection with pathogens. WASH is detected at attachment sites of Salmonella, and is involved in the entry of this pathogenic bacterium into the cell. Similarly, VPEF downregulation impairs the entry of Vaccinia virus. A surprising result, however, was that entry of Vaccinia is also blocked by extracellular antibodies directed against VPEF or by soluble VPEF in the culture medium, as if VPEF played the role of an extracellular receptor for the virus. These observations remain to be explained, because the WASH complex is cytosolic and no transmembrane complex is involved. The WASH subunit activates the Arp2/3 complex through its VCA domain at its C-terminus. This domain is about 100 amino-acid long. WASH stands for Wiskott-Aldrich Syndrome Protein and SCAR Homolog. This descriptive name has imposed itself and replaced the previously used Orf19. The organization of WASH is indeed very similar to the ones of WASP and SCAR/WAVE activators of the Arp2/3 complex. Based on this analogy, the N-terminal WAHD domain covering the first 300 amino-acids, is likely to be integral to the multiprotein complex similar to the SCAR/WAVE N-terminal domain in the pentameric SCAR/WAVE complex and to the N-terminal domain of N-WASP, which complexes N-WASP with WIP family proteins. The rest of WASH, including the VCA domain, is predicted to be unstructured. The VPEF subunit is of major interest, but confusing. VPEF was first characterized as a factor required for Vaccinia virus entry into mammalian cells, hence its name Vaccinia Penetration Factor. Unfortunately, this publication has been overlooked in the two recent publications reporting VPEF association with WASH. We used the name of the cDNA, KIAA0592, to refer to this protein. Gomez and Billadeau used the name FAM21, standing for FAMilly number 21 coined in automatic annotations of protein families. We recommend to use the name VPEF, which is anterior and descriptive. The major pool of the WASH complex and hence of VPEF is associated with endosomes. Nonetheless, the WASH complex is recruited to the plasma membrane upon infection with pathogens. WASH is detected at attachment sites of Salmonella, and is involved in the entry of this pathogenic bacterium into the cell. Similarly, VPEF downregulation impairs the entry of Vaccinia virus. A surprising result, however, was that entry of Vaccinia is also blocked by extracellular antibodies directed against VPEF or by soluble VPEF in the culture medium, as if VPEF played the role of an extracellular receptor for the virus. These observations remain to be explained, because the WASH complex is cytosolic and no transmembrane

| Name alternative | VPEF (KIAA0592, FAM21) | KIAA1033 | Strumpellin (KIAA0196) | WASH (Orf19) | CapZ α | CapZ β | Cdc53 |
|------------------|------------------------|----------|------------------------|---------------|--------|--------|------|
| M. musculus      | 100                    | 100      | 100                    | 100           | 100 (3) | 100    | 100  |
| H. sapiens       | 72.5 (3)               | 96.8     | 95.6                   | 84.2 (12-20)  | 96.5 (3) | 90.2   | 89.6 |
| X. tropicalis    | 43.9                   | 82.7     | 87.6                   | 69.3          | 88.1 (2)| 88.8   | 68.1 |
| D. rerio         | 37.9                   | 83.8     | 87.6                   | 67            | 79.3 (2)| 87.7   | 61.8 |
| D. melanogaster  | ND                     | 25.1     | 42.3                   | 27.9          | 61.4   | 77.6   | 31.7 |
| C. elegans       | ND                     | 17.5     | 19.2                   | 23.6          | 52.3   | 64.7   | 22.8 |
| Capitella sp     | ND                     | 61.9     | 63.4                   | 43.9          | 56.5   | 76.4   | 47.8 |
| L. gigantea      | ND                     | 60.6     | 66.6                   | 42.8          | 61.1   | 80.1   | 44.5 |
| M. brevicollis   | ND                     | 42.8     | 52.1                   | 33.3          | 54.5   | 70.1   | 33.1 |
| S. cerevisiae    | ND                     | ND       | ND                     | ND            | ND     | 25.8   | 44.4 |
| S. pombe         | ND                     | ND       | ND                     | ND            | 25     | 46.8   | ND   |
| D. discoideum    | ND                     | 41.1     | 48.6                   | 26            | 35.1   | 57.1   | ND   |
domain is predicted in the VPEF protein. VPEF is overall poorly conserved, and the percentage of identity drops so rapidly in vertebrates that invertebrate orthologs are not detected (Fig. 1). The N-terminal 350 amino-acids of VPEF are necessary and sufficient for building the WASH complex.7 This result is in line with the conservation concentrated in the N-terminal domain (Fig. 2). Right after this domain, VPEF is predicted to be disordered with high probability (Suppl. Fig. 2). This lack of intrinsic structure for the most part of the protein might explain why VPEF is highly sensitive to proteases and does not migrate at its expected size after purification of the WASH complex.1

The Strumpellin subunit is mutated in patients affected by hereditary spastic paraplegia. It is named after the physician Strümpell, who first described the disease more than 100 years ago.9 Three missense mutations have been identified in unrelated families.10 Strumpellin is one of the most conserved protein of the WASH complex, and these mutations are likely to affect the structure of Strumpellin. The mutant alleles of Strumpellin are dominant, suggesting that half a dose of WT Strumpellin is not sufficient to maintain a WT phenotype or that mutant Strumpellin blocks the assembly of functional WASH complexes despite the presence of WT Strumpellin in a dominant-negative manner.

The KIAA1033 subunit is also well conserved, similarly to Strumpellin. The last 100 amino-acids, however, correspond to a poorly conserved region, which is predicted to be intrinsically unstructured (Suppl. Figs. 2 and 4).

The Ccdc53 subunit is the second less conserved subunit after VPEF. It cannot be detected in D. discoideum, even though a functional ortholog is likely to exist in this species, where five out of seven subunits are detected (Fig. 1). Its name comes from Coiled-coil domain containing protein 53, because of a short and conserved coiled-coil predicted in the N-terminal domain. In this protein, two blocks of conserved residues alternate with two variable regions, predicted to be highly disordered (Suppl. Fig. 2).

Understanding how these seven subunits are assembled into a functional actin polymerization machine is of major fundamental importance and might also provide an explanation for why hereditary spastic paraplegia arises in patients affected by Strumpellin mutations.

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Note

Supplementary materials can be found at: www.landesbioscience.com/supplement/DeriveryCIB3-3-Sup.pdf

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