Supporting information 3 Figs: Supporting Phylogenetic analysis

Phylogenetic analysis methods are the same as described in the main manuscript with the exception that sequence alignment input in PhyML contained gaps.

Maximum likelihood phylogenetic analysis was performed in PhyML v3.0 server. Using the protein evolutionary model LG +G +F (previously calculated in Protest) and the number of bootstrap replicates was set to 1000. The resulting tree was visualized in Fig Tree V1.3.1 available at http://tree.bio.ed.ac.uk/software/figtree/ and rooted DME and CIN.
Bayesian phylogenetic analysis was performed using MrBayes v3.2.3 available in CIPRES Science Gateway V3.3. MrBayes was run for 1 million generations with the following parameters: rate matrix for aa=mixed, nruns=2, nchains=4, temp=0.2, sampling set to 1000 and burnin to 0.25. The resulting tree was visualized in Fig Tree V1.3.1 available at http://tree.bio.ed.ac.uk/software/figtree/ and rooted DME and CIN.

Phylogenetic analysis conducted in MEGA6 using Neighbor-Joining method. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (100 replicates) are shown next to the branches. The evolutionary distances were computed using the JTT matrix-based method and are in the units of the number of amino acid substitutions per site.