

Additional file 3: Bioinformatic analysis of proteins extracted from cell walls of *C. albicans* [20].

Data are from [20]. Proteins were extracted from *C. albicans* cell walls as described in Figure 3. Four successive extractions were performed : step 1 using SDS-DTT, step 2 using NaOH, step 3 using a β -1,3-glucanase and step 4 using an exochitinase. All proteins sequences were analyzed with bioinformatic programs to predict their sub-cellular localization. Proteins for which predictions by different bioinformatic programs are in conflict are classified as "not clear".

step 1: SDS-DTT extract	Predicted subcellular localization	Accession (a)	PSORT (b)	TargetP (c)	Predicted signal peptide (d)
	outside	CA1541 Phrp *	outside (0.370)	secretory pathway (0.946)	1-18
intracellular	CA0362	cytoplasm (0.650) endoplasmic reticulum (0.910) (C-term HDEL)	other (0.911)	1-29 or 1-32	
	CA0915	mitochondry (0.541)	secretory pathway (0.977)		
	CA1015	nucleus (0.760)	other (0.694), mitochondry (0.271)		
	CA1230	mitochondry (0.360)	other (0.843)		
	CA1691	endoplasmic reticulum (0.910) (C-term HDEL)	other (0.789), mitochondry (0.227)		
	CA1755	microbody (0.300)	secretory pathway (0.955)	1-23	
	CA2474	nucleus (0.760)	other (0.798)		
	CA2857	nucleus (0.980)	other (0.842)		
	CA3081	cytoplasm (0.650)	other (0.885)		
	CA3208	mitochondry (0.360)	mitochondry (0.663)		
	CA3483	cytoplasm (0.450)	other (0.275), mitochondry (0.299)		
	CA3534	cytoplasm (0.450)	other (0.887)		
	CA3874	cytoplasm (0.450)	other (0.793)		
	CA4671	mitochondry (0.853)	other (0.834)		
	CA4765	endoplasmic reticulum (0.550)	mitochondry (0.716)		
	CA4844	mitochondry (0.360)	other (0.704)		
	CA4862	nucleus (0.600)	other (0.934)		
	CA4959	cytoplasm (0.450)	other (0.923)		
	CA5135	cytoplasm (0.450)	other (0.928)		
	CA5180	cytoplasm (0.450)	other (0.797)		
	CA5892	mitochondry (0.471)	mitochondry (0.637)		
	CA5950	microbody (0.800)	other (0.604)		
not predictable	CA2810	not present in the database			
step 2: NaOH extract	outside	Hsp150p/Pir2p*			
intracellular	CA1691	mitochondry (0.360)	other (0.789), mitochondry (0.227)		
	CA2474	microbody (0.300)	other (0.798)		
	CA3483	mitochondry (0.360)	other (0.275), mitochondry (0.299)		
	CA3874	cytoplasm (0.450)	other (0.793)		
	CA4765	mitochondry (0.853)	mitochondry (0.716)		
	CA5180	cytoplasm (0.450)	other (0.797)		
	CA5892	mitochondry (0.471)	mitochondry (0.637)		

step 3: β-1,3-glucanase extract	outside	putative β -1,3-glucanase* Hsp150p/Pir2p* Phrp *		
	intracellular	CA1691 CA3874 CA5892	mitochondry (0.360) cytoplasm (0.450) mitochondry (0.471)	other (0.789), mitochondry (0.227) other (0.793) mitochondry (0.637)

step 4: exochitinase extract	outside	Hsp150p/Pir2p* Phrp *		
	intracellular	CA1691 CA2474 CA3874 CA5180 CA5892	mitochondry (0.360) microbody (0.300) cytoplasm (0.450) cytoplasm (0.450) mitochondry (0.471)	other (0.789), mitochondry (0.227) other (0.798) other (0.793) other (0.797) mitochondry (0.637)

colour code: proteins found in at least 2 fractions
 proteins found in at least 2 fractions

* proteins that failed to be identified to already annotated *C. albicans* proteins

(a) <http://genolist.pasteur.fr/CandidaDB/>

(b) PSORT : <http://psort.nibb.ac.jp/form.html> [29]

(b) TargetP: <http://www.cbs.dtu.dk/services/TargetP/> [30]

(d) Two sizes are indicated when different signal peptides are predicted by PSORT and TargetP. The first one is predicted with PSORT.