


genotype	WT	WT	$\Delta ppz1,2$
FLAG-Rps31	—	+	+
FLAG-Rpl40B	—	+	+
α -FLAG			
α -G6PDH			

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FLAG-Rps31	—	+	+
FLAG-Rpl40B	—	+	+
α -FLAG			
α -G6PDH			

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FLAG-Rps31	—	+	+
FLAG-Rpl40B	—	+	+
α -FLAG			
α -G6PDH			

Figure 1, figure supplement 3. Quantification of cellular ubiquitin levels by analysis of slot blots. Yeast strains (SEY6210 background) with endogenously FLAG-tagged ubiquitin (at the *RPL40B* and *RPS31* loci with native promoter and terminator containing an N-terminal 3xFLAG tag) were grown to mid-log phase and precipitated in 10% cold TCA. Total cell lysates from solubilized pellets were bound to PVDF membranes using a vacuum manifold and immunoblotting analysis was performed as indicated. Three biological replicate experiments are shown.