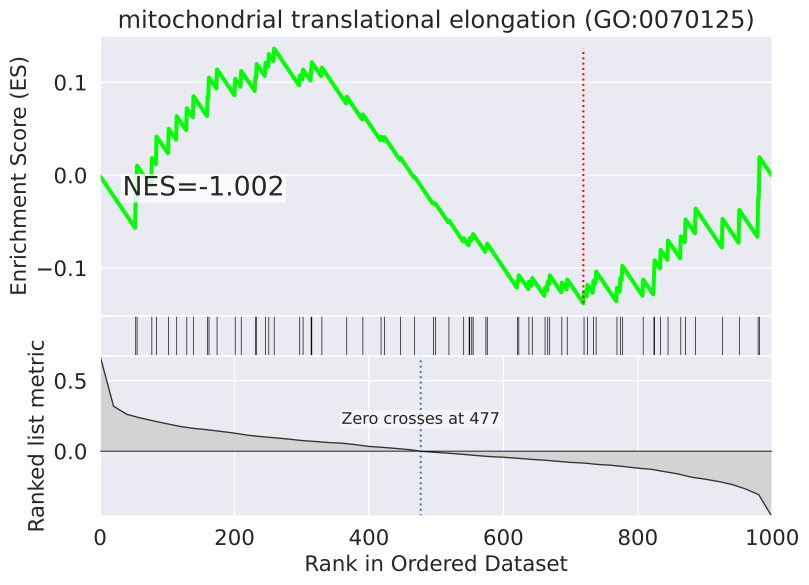
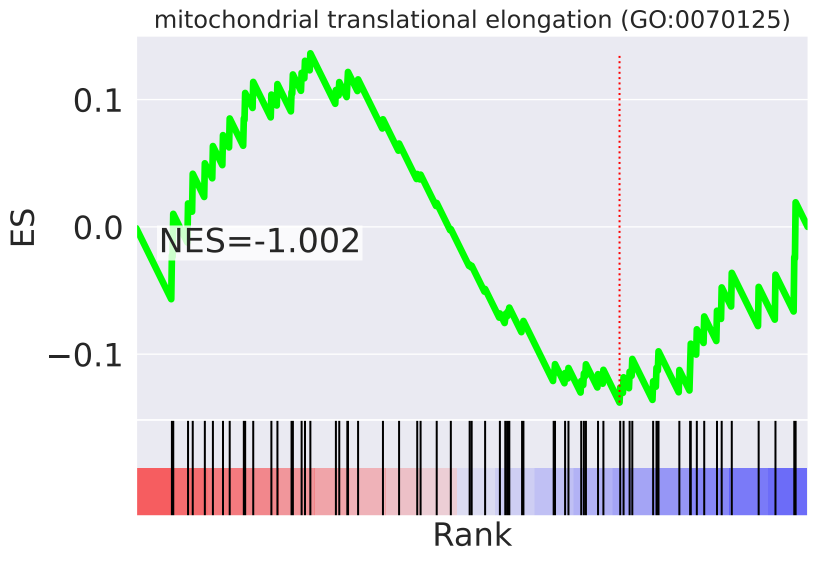
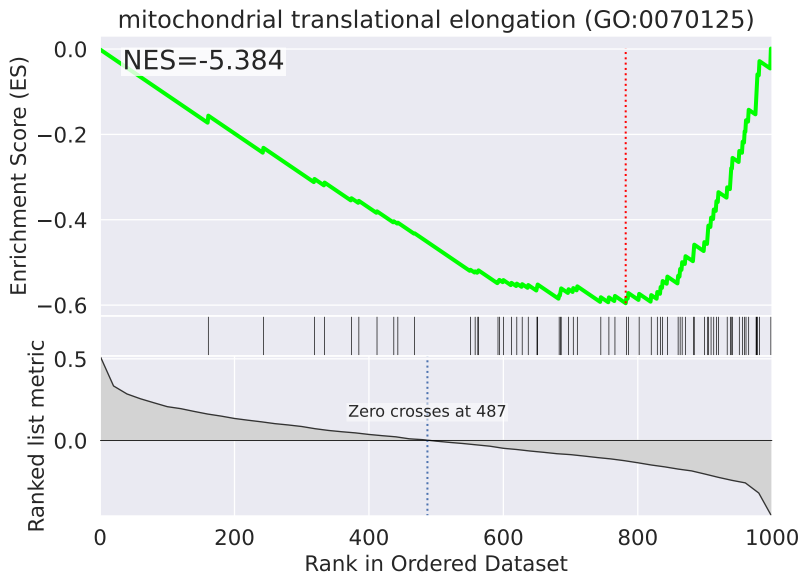


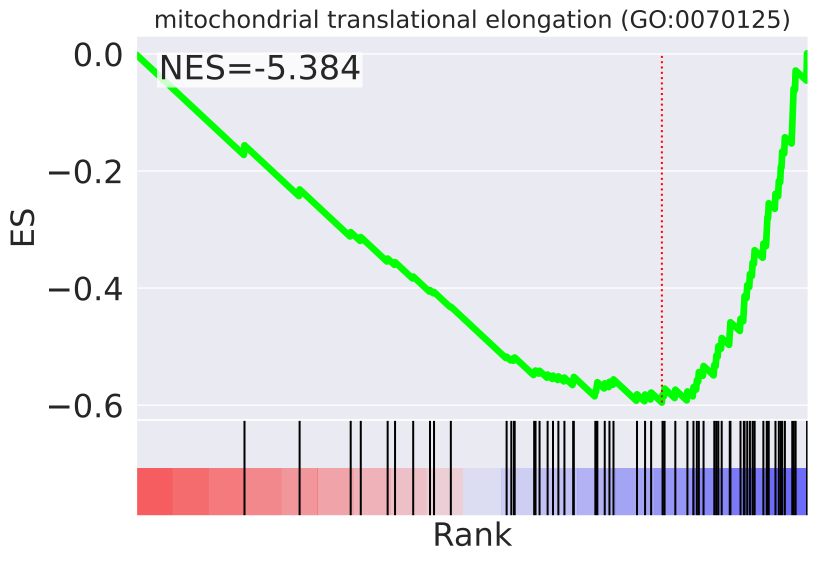
NES	SET
-5.237	mitochondrial translational elongation (GO:0070125)
-5.039	mitochondrial translational termination (GO:0070126)
-3.291	translation (GO:0006412)
-2.972	mitochondrial translation (GO:0032543)
2.677	protein ubiquitination involved in ubiquitin-dependent protein catabolic process (GO:0042787)
2.676	neutrophil degranulation (GO:0043312)
2.646	purine ribonucleoside monophosphate biosynthetic process (GO:0009168)
-2.374	protein homooligomerization (GO:0051260)
-2.368	negative regulation of transcription, DNA-templated (GO:0045892)
2.263	mitotic spindle organization (GO:0007052)
-2.253	tRNA aminoacylation for protein translation (GO:0006418)
2.227	transcription-coupled nucleotide-excision repair (GO:0006283)
2.196	mitochondrial electron transport, NADH to ubiquinone (GO:0006120)
-2.181	mitochondrial electron transport, cytochrome c to oxygen (GO:0006123)
2.128	negative regulation of telomere maintenance via telomerase (GO:0032211)



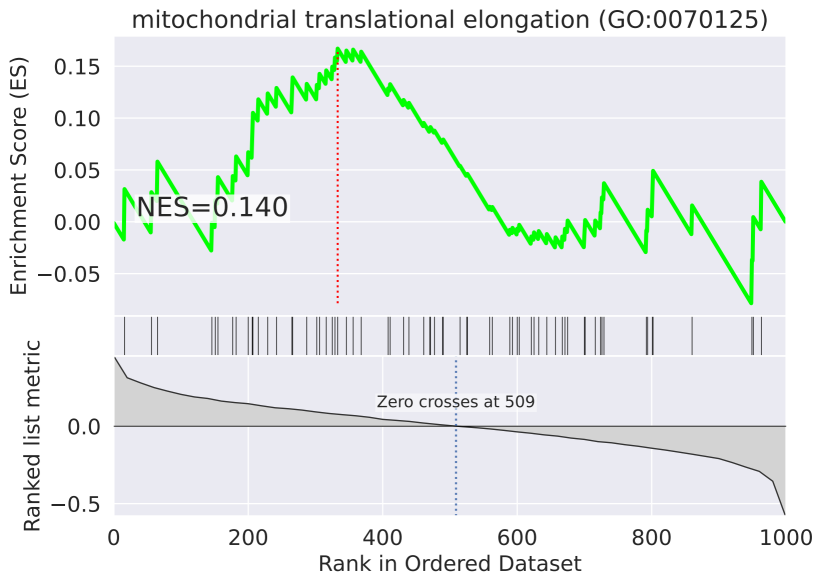


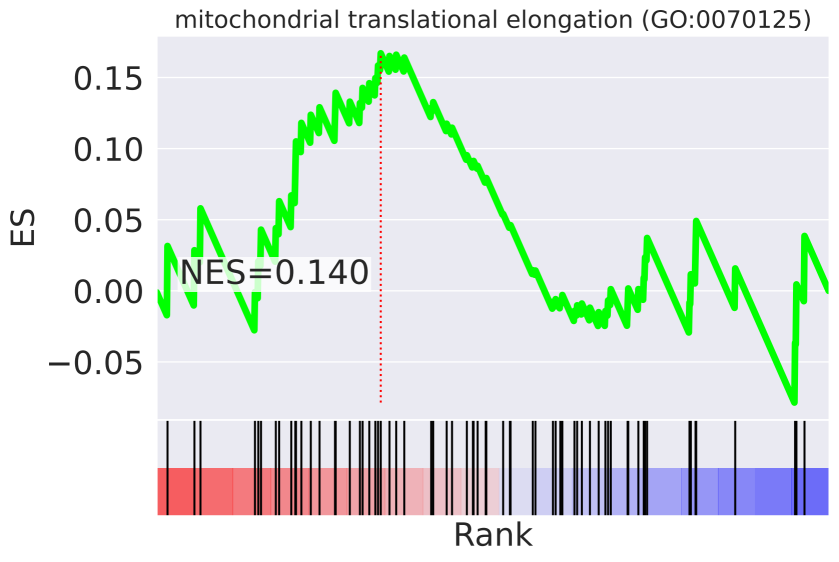
NES	SET
-3.279	purine ribonucleoside monophosphate biosynthetic process (GO:0009168)
-2.843	insulin receptor signaling pathway (GO:0008286)
-2.767	Ras protein signal transduction (GO:0007265)
2.710	regulation of mRNA stability (GO:0043488)
-2.486	regulation of cholesterol biosynthetic process (GO:0045540)
-2.464	axon guidance (GO:0007411)
-2.454	ERBB2 signaling pathway (GO:0038128)
-2.312	cholesterol biosynthetic process (GO:0006695)
-2.309	epidermal growth factor receptor signaling pathway (GO:0007173)
-2.300	leukocyte migration (GO:0050900)
-2.293	response to endoplasmic reticulum stress (GO:0034976)
2.213	regulation of apoptotic process (GO:0042981)
2.188	regulation of ubiquitin-protein ligase activity involved in mitotic cell cycle (GO:0051439)
-2.160	positive regulation of mitotic cell cycle (GO:0045931)
-2.135	fibroblast growth factor receptor signaling pathway (GO:0008543)



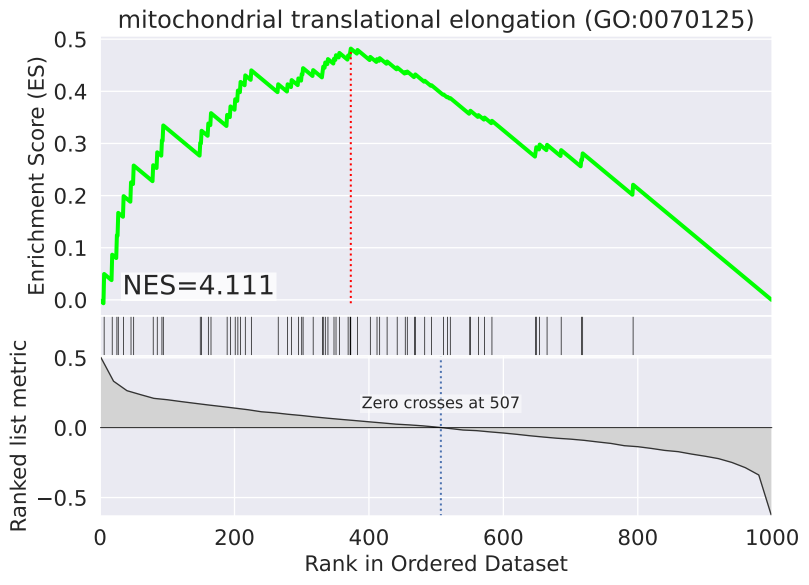


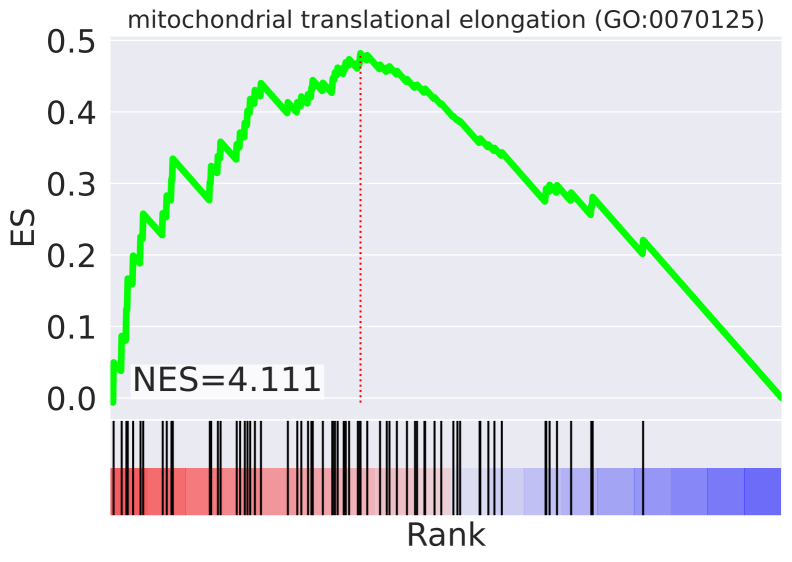
NES	SET
-5.384	mitochondrial translational elongation (GO:0070125)
-5.241	mitochondrial translational termination (GO:0070126)
-4.313	mitochondrial respiratory chain complex I assembly (GO:0032981)
-4.127	mitochondrial electron transport, NADH to ubiquinone (GO:0006120)
3.475	positive regulation of cell proliferation (GO:0008284)
-3.259	mitochondrial translation (GO:0032543)
3.103	negative regulation of apoptotic process (GO:0043066)
3.033	MAPK cascade (GO:0000165)
2.968	positive regulation of cell growth (GO:0030307)
2.896	Fc-gamma receptor signaling pathway involved in phagocytosis (GO:0038096)
2.860	Ras protein signal transduction (GO:0007265)
2.834	retrograde vesicle-mediated transport, Golgi to ER (GO:0006890)
2.782	positive regulation of protein phosphorylation (GO:0001934)
-2.780	translation (GO:0006412)
2.760	endosomal transport (GO:0016197)



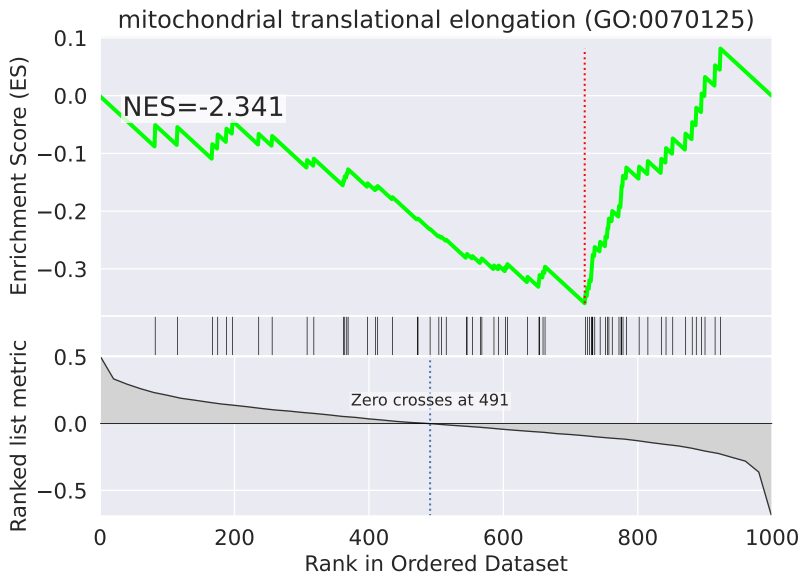


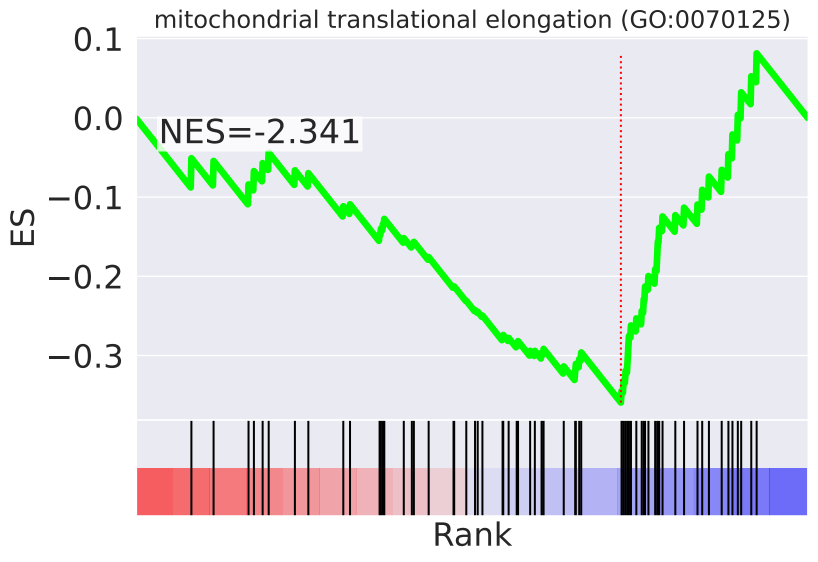
NES	SET
-3.332	membrane organization (GO:0061024)
2.962	protein ubiquitination involved in ubiquitin-dependent protein catabolic process (GO:0042787)
2.839	mitotic cell cycle (GO:0000278)
2.735	mitotic cytokinesis (GO:0000281)
2.721	mitotic metaphase plate congression (GO:0007080)
-2.661	purine ribonucleoside monophosphate biosynthetic process (GO:0009168)
2.620	anaphase-promoting complex-dependent catabolic process (GO:0031145)
-2.429	regulation of cholesterol biosynthetic process (GO:0045540)
-2.388	protein autophosphorylation (GO:0046777)
2.366	mRNA splicing, via spliceosome (GO:0000398)
2.303	chromosome segregation (GO:0007059)
2.251	DNA replication (GO:0006260)
2.238	intrinsic apoptotic signaling pathway (GO:0097193)
2.209	tRNA modification (GO:0006400)
2.196	sister chromatid cohesion (GO:0007062)



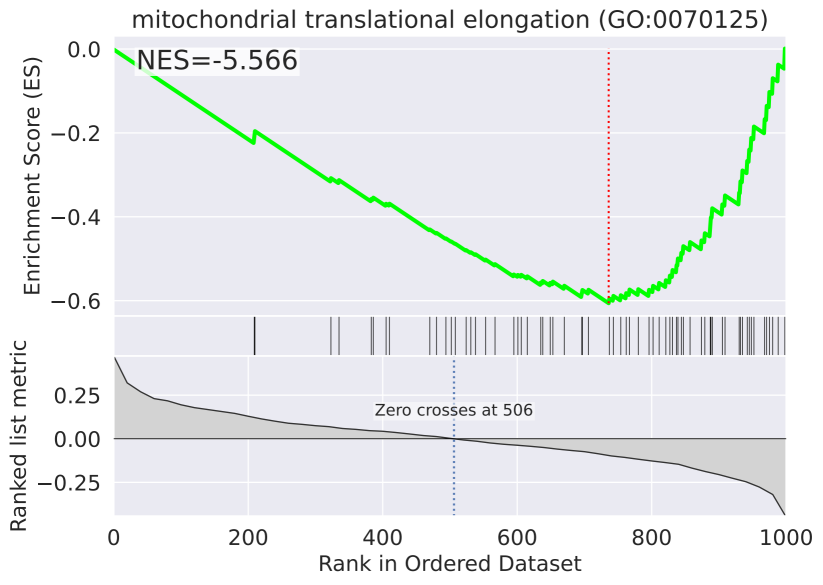


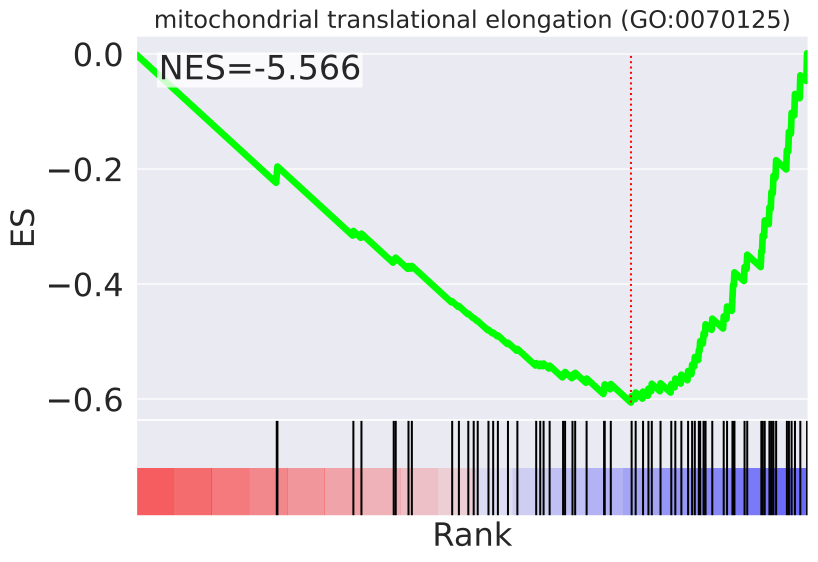
NES	SET
4.189	mitochondrial translational termination (GO:0070126)
4.111	mitochondrial translational elongation (GO:0070125)
-3.523	positive regulation of cell proliferation (GO:0008284)
3.130	mitochondrial respiratory chain complex I assembly (GO:0032981)
3.068	mitochondrial electron transport, NADH to ubiquinone (GO:0006120)
2.897	protein deubiquitination (GO:0016579)
-2.759	protein autophosphorylation (GO:0046777)
-2.749	signal transduction (GO:0007165)
-2.698	positive regulation of cell migration (GO:0030335)
-2.617	epidermal growth factor receptor signaling pathway (GO:0007173)
-2.578	nuclear-transcribed mRNA poly(A) tail shortening (GO:0000289)
-2.557	movement of cell or subcellular component (GO:0006928)
-2.556	Fc-gamma receptor signaling pathway involved in phagocytosis (GO:0038096)
-2.493	protein K63-linked ubiquitination (GO:0070534)
-2.465	vascular endothelial growth factor receptor signaling pathway (GO:0048010)



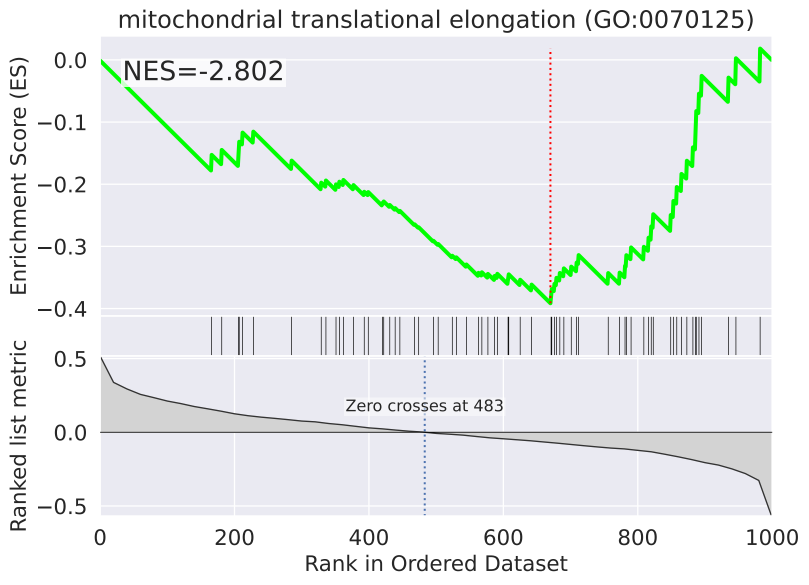


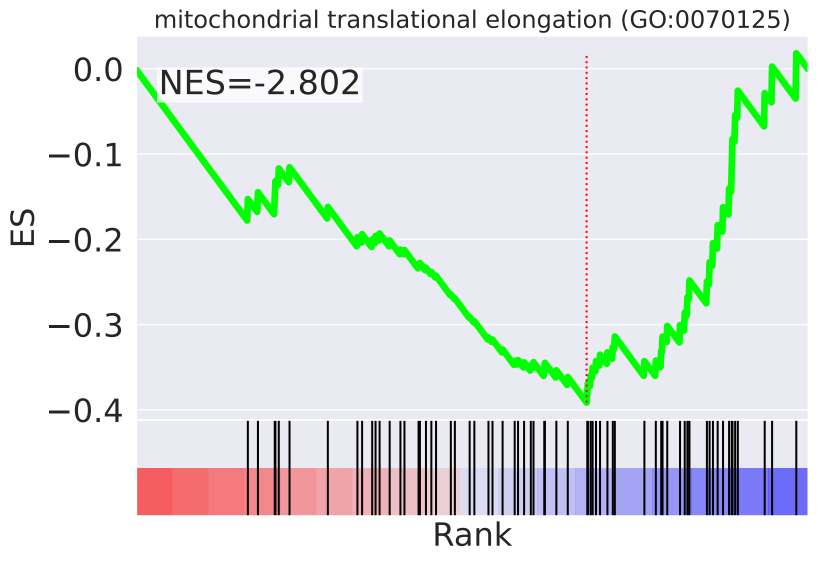
NES	SET
-3.442	tricarboxylic acid cycle (GO:0006099)
-3.180	purine ribonucleoside monophosphate biosynthetic process (GO:0009168)
2.613	DNA repair (GO:0006281)
2.601	integrin-mediated signaling pathway (GO:0007229)
2.466	positive regulation of sequence-specific DNA binding transcription factor activity (GO:0051091)
-2.425	cellular nitrogen compound metabolic process (GO:0034641)
-2.346	membrane organization (GO:0061024)
-2.341	mitochondrial translational elongation (GO:0070125)
-2.310	androgen receptor signaling pathway (GO:0030521)
-2.284	retrograde transport, endosome to Golgi (GO:0042147)
-2.204	tRNA aminoacylation for protein translation (GO:0006418)
2.185	protein K63-linked ubiquitination (GO:0070534)
2.167	vascular endothelial growth factor receptor signaling pathway (GO:0048010)
2.152	transforming growth factor beta receptor signaling pathway (GO:0007179)
2.151	protein autoubiquitination (GO:0051865)



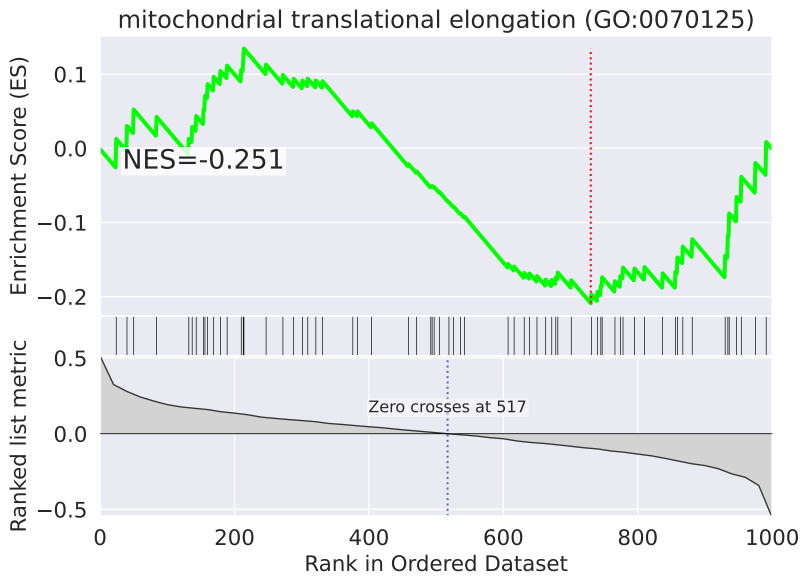


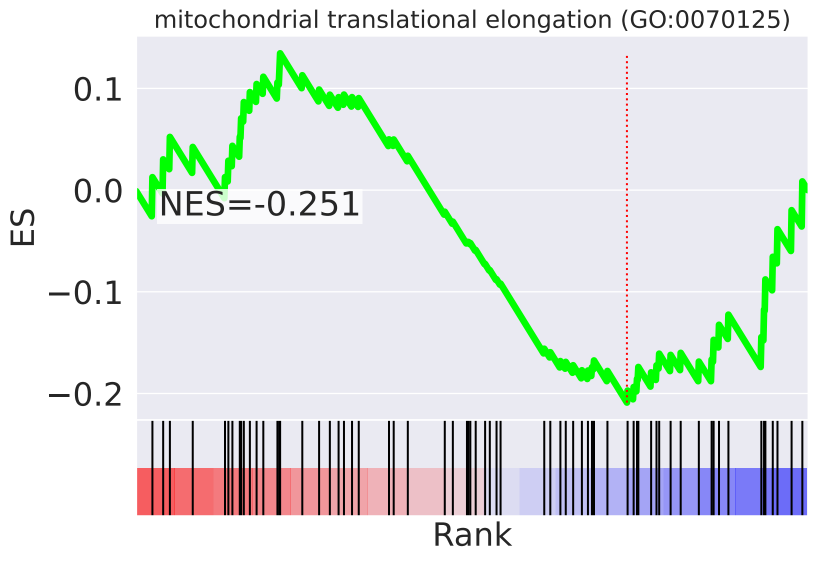
NES	SET
-5.744	mitochondrial translational termination (GO:0070126)
-5.566	mitochondrial translational elongation (GO:0070125)
-4.183	mitochondrial respiratory chain complex I assembly (GO:0032981)
-3.861	mitochondrial electron transport, NADH to ubiquinone (GO:0006120)
2.765	sister chromatid cohesion (GO:0007062)
2.562	purine ribonucleoside monophosphate biosynthetic process (GO:0009168)
2.486	Fc-gamma receptor signaling pathway involved in phagocytosis (GO:0038096)
-2.421	nucleosome disassembly (GO:0006337)
-2.420	regulation of macroautophagy (GO:0016241)
2.382	regulation of cellular response to heat (GO:1900034)
-2.353	translation (GO:0006412)
2.327	axon guidance (GO:0007411)
2.325	cytokinesis (GO:0000910)
2.252	epidermal growth factor receptor signaling pathway (GO:0007173)
-2.235	endosomal transport (GO:0016197)





NES	SET
-3.194	regulation of cholesterol biosynthetic process (GO:0045540)
3.068	mitotic nuclear envelope disassembly (GO:0007077)
3.010	rRNA processing (GO:0006364)
-2.955	histone H3 acetylation (GO:0043966)
2.876	mitotic cell cycle (GO:0000278)
-2.802	mitochondrial translational elongation (GO:0070125)
2.703	protein sumoylation (GO:0016925)
2.687	double-strand break repair (GO:0006302)
2.648	platelet aggregation (GO:0070527)
2.645	tRNA export from nucleus (GO:0006409)
2.645	regulation of glucose transport (GO:0010827)
2.645	regulation of gene silencing by miRNA (GO:0060964)
2.628	telomere capping (GO:0016233)
2.611	mitotic metaphase plate congression (GO:0007080)
2.602	intracellular transport of virus (GO:0075733)
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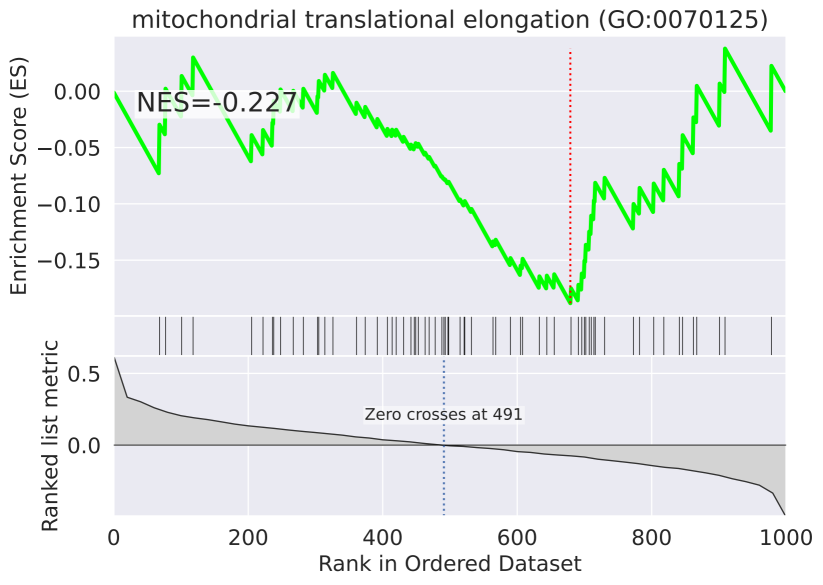


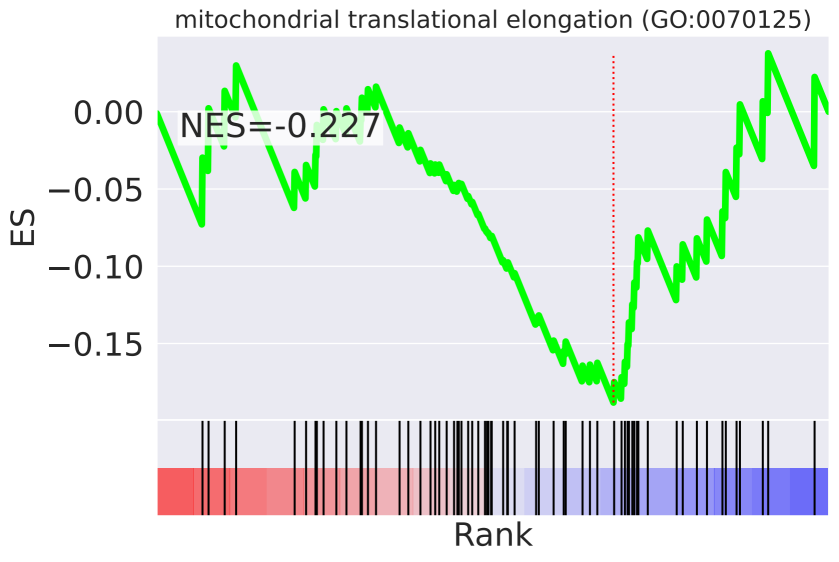


double-strand break repair via nonhomologous end joining (GO:0006303)
cell differentiation (GO:0030154)
anaphase-promoting complex-dependent catabolic process (GO:0031145)
mitochondrial respiratory chain complex I assembly (GO:0032981)
post-translational protein modification (GO:0043687)
nucleus organization (GO:0006997)
cholesterol biosynthetic process (GO:0006695)
telomere capping (GO:0016233)
protein K48-linked ubiquitination (GO:0070936)
negative regulation of canonical Wnt signaling pathway (GO:0090090)
spermatogenesis (GO:0007283)
protein K11-linked ubiquitination (GO:0070979)
histone H4 acetylation (GO:0043967)
viral life cycle (GO:0019058)
positive regulation of sequence-specific DNA binding transcription factor activity (GO:0051091)

NES

SET





NES	SET
-3.213	regulation of cholesterol biosynthetic process (GO:0045540)
3.121	vascular endothelial growth factor receptor signaling pathway (GO:0048010)
3.106	cell-matrix adhesion (GO:0007160)
2.990	integrin-mediated signaling pathway (GO:0007229)
2.944	intracellular signal transduction (GO:0035556)
-2.911	cellular nitrogen compound metabolic process (GO:0034641)
2.767	Fc-gamma receptor signaling pathway involved in phagocytosis (GO:0038096)
2.685	ubiquitin-dependent protein catabolic process (GO:0006511)
2.656	protein phosphorylation (GO:0006468)
2.647	retrograde vesicle-mediated transport, Golgi to ER (GO:0006890)
2.596	platelet aggregation (GO:0070527)
2.506	movement of cell or subcellular component (GO:0006928)
2.440	ubiquitin-dependent ERAD pathway (GO:0030433)
2.440	retrograde protein transport, ER to cytosol (GO:0030970)
-2.435	tricarboxylic acid cycle (GO:0006099)