

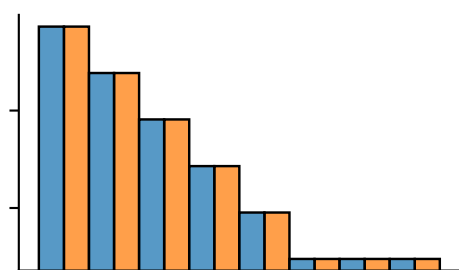
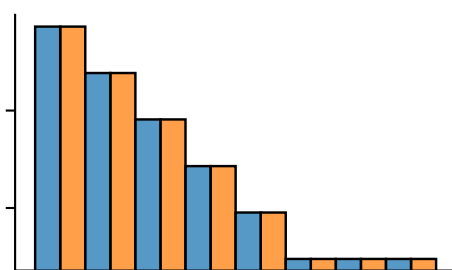
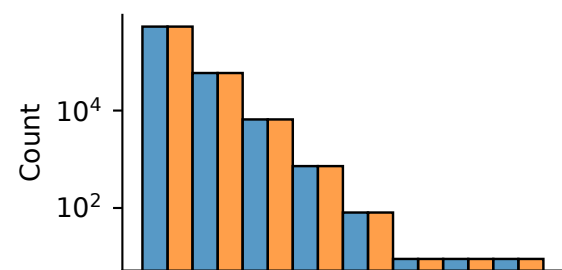
n=256 agents per node

normomutator hypermutator

available beneficial mutations = 1

available beneficial mutations = 2

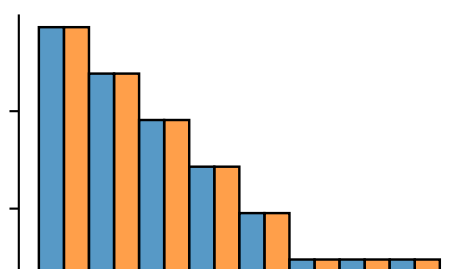
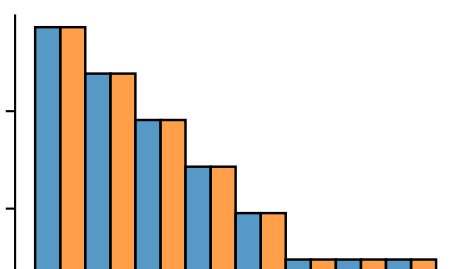
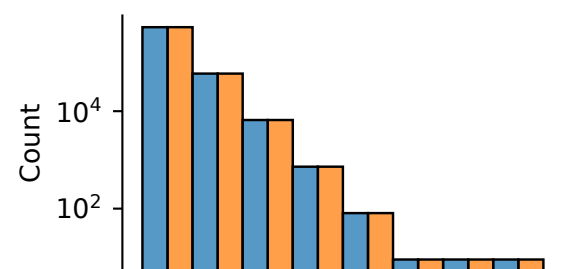
available beneficial mutations = 3



available beneficial mutations = 4

available beneficial mutations = 6

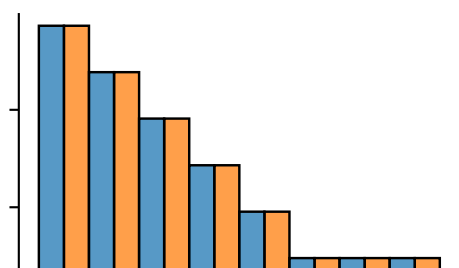
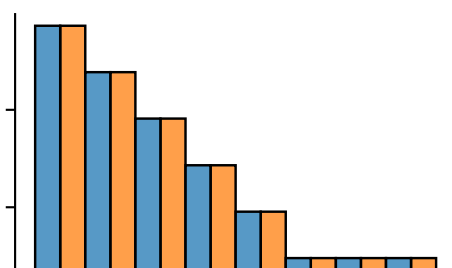
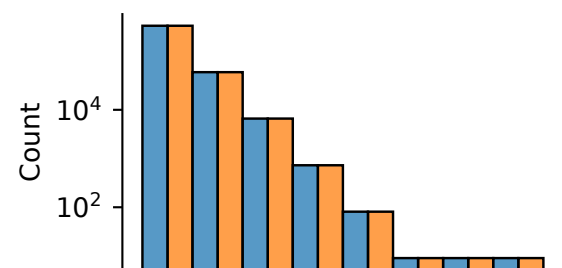
available beneficial mutations = 8



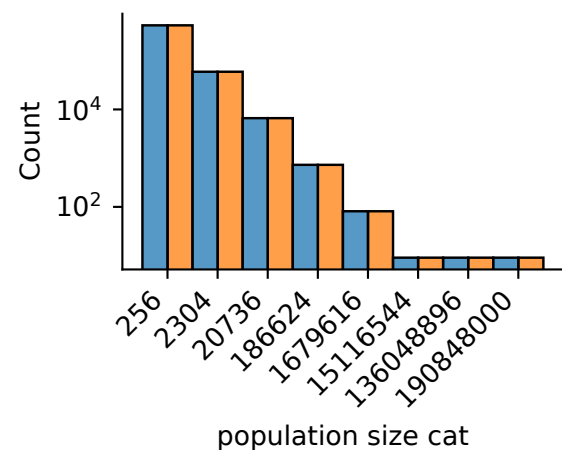
available beneficial mutations = 10

available beneficial mutations = 12

available beneficial mutations = 14



available beneficial mutations = 16



256 2304 20736 186624 1679616 15116544 136048896 190848000
population size cat

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population size cat