Table1. The time saving results for CCSD iterations of ground state energy calculation in double- and mixed-precision are shown in the tables below. The calculations were running for water clusters consisting of 1 to 7 water molecules with 3-21g basis set. It is shown that the proposed mixed-precision method has comparable accuracy as double-precision calculation and can reach the expected time saving with the increasing system size. (\*Accuracy column shows how many digits after the decimal places can the results match with PSI4 package.)

Number of H <sub>2</sub> O	Double-precision(dp)			Mixed-precision(mp)			
	Time/s	Iterations	Accuracy*	Time /s	Iterations	Accuracy	Time savings
1	0.19	17	6	0.16	17	6	0.84210
2	2.69	15	6	1.64	15	6	0.60966
3	23.28	16	6	13.14	16	6	0.564433
4	122.95	16	6	61.97	15	5	0.50402
5	428.63	17	6	239.43	17	6	0.55859
6	1211.93	17	6	617.63	16	6	0.50962
7	2790.41	17	6	1497.20	17	6	0.536552