

IM Calculation Refactor

Please read the readme @ https://github.com/ucgmsim/IM_calculation/blob/master/README.md for instructions on how to run the code.

DONE

- IM calculations have been separated from the "old post-processing" repository, extracting the relevant functions and classes.
- IM values validated on Hypocentre and Kupe against "old post-processing" on the same data.
- Two types of workflows: text based (most likely observations) and binary based (simulations once the binary workflow is in place).
 - If binary workflow does not happen soon, the text based option will be used in both cases.
- Outputs match the formats requested in [File Formats Used In Ground Motion Simulation](#) and should therefore be usable on the upcoming Non-ergodic codes.
- Tested on very simple multi-process on Kupe with good speed-up using 40 and 80 cores. For the sample, 2228 stations were used

Machine	Cores	Time
Hypocentre	1	132m
Hypocentre	8	8.7m
Kupe	40	27m
Kupe	80	

OUTPUT STRUCTURE

With command : `python calculate_ims.py ../BB.bin b -o /home/yzh231/ -i Albury_666_999 -r Albury -t s -v 18p3 -n 112A -m PGV pSA -p 0.02 0.03 -e -c geom -np 2`

- input file path:** `../BB.bin`
- b:** input file type is **binary**
- o:** output result csvs location is `/home/yzh231`, default is `/home/$user`
- i:** unique identifier/runname of the simrun and output folder name are **Albury_666_999**, default is `'all_station_ims'`. This attribute will be stored in the meta data file.
- r:** rupture name is **Albury**, default is unknown. This attribute will be stored in the meta data file.
- t:** type of simrun is **simulated**, default is unknown. This attribute will be stored in the meta data file.
- v:** version of simrun is **v18p3**, default is **XXpY**. This attribute will be stored in the meta data file.
- n:** station names used to perform im claculation are **112A**, default is all the stations in the binary file
- m:** measures used to perform im calculation are **PGV and pSA**, default is all the measures
- p:** period of pSA used to perform im calculation are **0.02 0.03**, default is Karim's 15 periods
- e:** In addition to the period specified by `-p` option, **use extended 100 period of pSA**, default not using
- c:** component of waveform acceleration used to perform im calculation is **geom**, default is `'090, 000, ver'`
- np:** number of processors used to perform im calculation is **2**, default is 2

The result is outputted to the following location, where:

- `'Albury_666_999'` is the folder that contains all outputs. The folder name `'Albury_666_999'` is made of the string specified by the `'-i'` argument. Default is `'all_station_ims'` if not specified.
- `'Albury_666_999.csv'` is the summary csv file that contains all stations' im calculations. The summary file name is made of the string specified by the `'-i'` argument.
- `'Albury_666_999_imcalc.info'` is the meta data file. The meta data file name is made of the string specified by the `'-i'` argument.
- `'station'` is the folder that contains all individual station's im calculations. The folder name is defaulted and cannot be specified by the user.
- `'112A_geom.csv'` is the individual csv file that contains geom component im calculation for station 112A. Each name of the individual station csv file name is made of `station_name + component`

```
yzh231@hypocentre ~ % tree Albury_666_999
Albury_666_999
├── Albury_666_999.csv
├── Albury_666_999_imcalc.info
├── stations
│   └── 112A_geom.csv
```

TEST FOR CALCUALTE_IMS.PY

All the steps below are to be carried out in hypocentre

1.Generate summary benchmark:

The following steps should only be performed once for each selected binary file

1. Select a source binary file: `/nesi/transit/nesi00213/RunFolder/daniel.lagrava/Kelly_VMSI_Kelly-h0p4_EMODv3p0p4_180531/BB/Cant1D_v2-midQ_leer_hfnp2mm+_rvf0p8_sd50_k0p045/Kelly_HYP01-03_S1244/Acc/BB_with_siteamp.bin`
2. Identify corresponding database for the selected source binary file: `/home/nesi00213/RunFolder/wd16/database_old_pp/database.db`
3. Find the script to extract benchmark im value files from the database in step 2: `/nesi/projects/nesi00213/dev/impp_datasets/extract_ims.sql`
4. Create a folder to store benchmark files. eg `benchmark_im_sims`
5. Execute `extract_ims.sql` in database.db 4 times with specified components. eg: 'ver'

```

1 select station_name, 'ver' as component,
2 (SELECT value FROM intensity_measure WHERE intensity_measure.station_id = stations.id AND measure_name = 'PGA' AND component = 'ver') as PGA,
3 (SELECT value FROM intensity_measure WHERE intensity_measure.station_id = stations.id AND measure_name = 'PGV' AND component = 'ver') as PGV,
4 (SELECT value FROM intensity_measure WHERE intensity_measure.station_id = stations.id AND measure_name = 'CAV' AND component = 'ver') as CAV,
5 (SELECT value FROM intensity_measure WHERE intensity_measure.station_id = stations.id AND measure_name = 'AI' AND component = 'ver') as AI,
6 (SELECT value FROM intensity_measure WHERE intensity_measure.station_id = stations.id AND measure_name = 'Ds575' AND component = 'ver') as Ds575,
7 (SELECT value FROM intensity_measure WHERE intensity_measure.station_id = stations.id AND measure_name = 'Ds595' AND component = 'ver') as Ds595,
8 (SELECT value FROM intensity_measure WHERE intensity_measure.station_id = stations.id AND measure_name = 'MMT' AND component = 'ver') as MMT,
9 (SELECT value FROM intensity_measure WHERE intensity_measure.station_id = stations.id AND measure_name = 'pSA' AND period < 0.01 * 1.001 AND period > 0.01 * 0.999 AND component = 'ver') as pSA_0,
10 (SELECT value FROM intensity_measure WHERE intensity_measure.station_id = stations.id AND measure_name = 'pSA' AND period < 0.0107226722201032 * 1.001 AND period > 0.0107226722201032 * 0.999
11 (SELECT value FROM intensity_measure WHERE intensity_measure.station_id = stations.id AND measure_name = 'pSA' AND period < 0.0114975699539774 * 1.001 AND period > 0.0114975699539774 * 0.999
12 (SELECT value FROM intensity_measure WHERE intensity_measure.station_id = stations.id AND measure_name = 'pSA' AND period < 0.0123284673944207 * 1.001 AND period > 0.0123284673944207 * 0.999
13 (SELECT value FROM intensity_measure WHERE intensity_measure.station_id = stations.id AND measure_name = 'pSA' AND period < 0.0132194114846603 * 1.001 AND period > 0.0132194114846603 * 0.999
14 (SELECT value FROM intensity_measure WHERE intensity_measure.station_id = stations.id AND measure_name = 'pSA' AND period < 0.0141747416292681 * 1.001 AND period > 0.0141747416292681 * 0.999
15 (SELECT value FROM intensity_measure WHERE intensity_measure.station_id = stations.id AND measure_name = 'pSA' AND period < 0.0151991108295293 * 1.001 AND period > 0.0151991108295293 * 0.999
16 (SELECT value FROM intensity_measure WHERE intensity_measure.station_id = stations.id AND measure_name = 'pSA' AND period < 0.0162975083462064 * 1.001 AND period > 0.0162975083462064 * 0.999
17 (SELECT value FROM intensity_measure WHERE intensity_measure.station_id = stations.id AND measure_name = 'pSA' AND period < 0.017475284000768 * 1.001 AND period > 0.017475284000768 * 0.999
18 (SELECT value FROM intensity_measure WHERE intensity_measure.station_id = stations.id AND measure_name = 'pSA' AND period < 0.0187381742286038 * 1.001 AND period > 0.0187381742286038 * 0.999

```

6. Export results to `benchmark_im_sims/benchmark_im_sim_ver.csv`. Click OK and don't change anything when 'Export data as csv' window prompts

station_name	component	PGA	PGV	CAV	AI	Ds575	Ds595	MMT	pSA_0p01	pSA_0p0107226722201032	pSA_0p0114975699539774	pSA_0p0123284673944207
2002199	ver	0.13105402603094	14.6578	0.25730606504577	11.4951405103342	3.855	11.13	6.57477738151169	0.131223057861578	0.131213055319727	0.13114649761158	0.13149840211
2000228	ver	0.02438328264896	4.83675	0.102252705927625	1.08737408488283	10.85	16.945	5.05318950343977	0.0245020293301113	0.0245139476883761	0.0245139476883761	0.024509006
2000224	ver	0.0389867482160999	5.73397	0.171694156945117	3.0522438390113	7.87	19.805	5.28671911581643	0.03907508850342	0.0391013684259	0.03910500611082	0.0391337837
41000af	ver	0.047866466293078	5.71101	0.202349174280887	4.17396864645412	9.855	17.045	5.281212891488	0.0479205514511365	0.0479027266039725	0.047905954953974	0.0478678709
52002359	ver	0.0553525993883902	9.2735	0.23500987591261	6.68968827817495	9.335	14.475	5.94649001711633	0.0555370515354537	0.0555694174641322	0.055613729248709	0.0556631104
1000187	ver	0.0593109072375246	8.50213	0.151733973433071	3.366192738885	6.41	13.22	5.82730766209323	0.0595141812766518	0.059529133016151	0.059518510752993	0.0596937274
72002356	ver	0.035626197573849	3.89133	0.120135541877813	1.50417854429293	8.38	18.335	4.57470987666496	0.0357095548144364	0.0357021564462068	0.0357217462613994	0.0357369871
820020a5	ver	0.041832823493309	3.40456	0.141207591250714	2.23761245837024	8.195	19.01	4.89321489961055	0.0419401053890726	0.0419568048675201	0.0420111344100175	0.04202369871
920020a6	ver	0.025231498470953	3.40659	0.116108725292778	1.38348919842566	9.435	18.635	4.60398678146469	0.0252891832668121	0.0252926831335556	0.02528546418741	0.0253098001
10200032b	ver	0.018793027321763	2.70051	0.084466961586555	0.70039323154614	11.54	19.025	4.41892598012615	0.0188118308450197	0.0188173629176038	0.018824940450606	0.0188301041

Export data as CSV

Column names in first line ☒

Field separator: ,

Quote character: "

Cancel OK

7. Repeat step 4 and 5 with different components: '090', '000', 'geom'
8. Now you have 4 summary benchmark files `benchmark_im_sim_090/000/ver/geom.csv`

2. Generate test input files

1. Follow the instruction in [Binary Workflow](#) FAQ, we can generate single waveform files. These waveforms are intended for the testing of ascii functionality of calculate_ims.py. Open a python cell

```
from qcore.timeseries import BBSeis
bb = BBSeis('nesi/transit/nesi00213/RunFolder/daniel.lagrava/Kelly_VMSI_Kelly-h0p4_EMODv3p0p4_180531/BB
/Cant1D_v2-midQ_leer_hfnp2mm+_rvf0p8_sd50_k0p045/Kelly_HYP01-03_S1244/Acc/BB_with_siteamp.bin')
bb.all2txt(self, prefix='/home/$user/benchmark_im_sim_waveforms/', f='acc'):
```

Now we have all the waveforms.

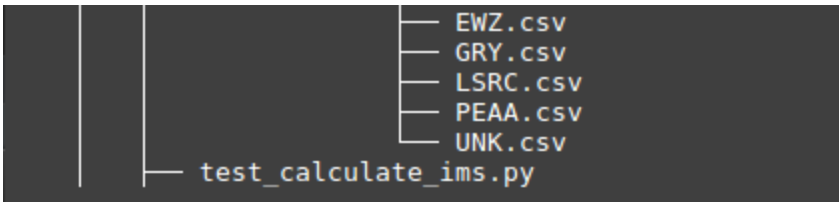
```
0002014.ver 0002307.000 0002540.090 000281f.ver 0002bce.000 10002b7.090 100086e.ver 1
0002015.000 0002307.090 0002540.ver 0002820.000 0002bce.090 10002b7.ver 1000871.000 1
0002015.090 0002307.ver 0002541.000 0002820.090 0002bce.ver 10002ca.000 1000871.090 1
0002015.ver 0002308.000 0002541.090 0002820.ver 0002bcf.000 10002ca.090 1000871.ver 1
0002016.000 0002308.090 0002541.ver 0002821.000 0002bcf.090 10002ca.ver 1000882.000 1
0002016.090 0002308.ver 00025e0.000 0002821.090 0002bcf.ver 10002cc.000 1000882.090 1
0002016.ver 0002309.000 00025e0.090 0002821.ver 0002bd0.000 10002cc.090 1000882.ver 1
0002017.000 0002309.090 00025e0.ver 0002822.000 0002bd0.090 10002cc.ver 10008cc.000 1
0002017.090 0002309.ver 00025e1.000 0002822.090 0002bd0.ver 10002d9.000 10008cc.090 1
0002017.ver 000230a.000 00025e1.090 0002822.ver 0002bd1.000 10002d9.090 10008cc.ver 1
0002018.000 000230a.090 00025e1.ver 0002823.000 0002bd1.090 10002d9.ver 10008cf.000 1
0002018.090 000230a.ver 00025e2.000 0002823.090 0002bd1.ver 10002e3.000 10008cf.090 1
0002018.ver 00023ab.000 00025e2.090 0002823.ver 0002c82.000 10002e3.090 10008cf.ver 1
00020c1.000 00023ab.090 00025e2.ver 0002824.000 0002c82.090 10002e3.ver 10008d3.000 1
00020c1.090 00023ab.ver 00025e3.000 0002824.090 0002c82.ver 10002f2.000 10008d3.090 1
00020c1.ver 00023ac.000 00025e3.090 0002824.ver 0002c83.000 10002f2.090 10008d3.ver 1
00020c2.000 00023ac.090 00025e3.ver 0002825.000 0002c83.090 10002f2.ver 10008de.000 1
00020c2.090 00023ac.ver 00025e4.000 0002825.090 0002c83.ver 10002ff.000 10008de.090 1
00020c2.ver 00023ad.000 00025e4.090 0002825.ver 0002c84.000 10002ff.090 10008de.ver 1
00020c3.000 00023ad.090 00025e4.ver 0002826.000 0002c84.090 10002ff.ver 10008e4.000 1
00020c3.090 00023ad.ver 00025e5.000 0002826.090 0002c84.ver 1000302.000 10008e4.090 1
00020c3.ver 00023ae.000 00025e5.090 0002826.ver 0002c85.000 1000302.090 10008e4.ver 1
00020c4.000 00023ae.090 00025e5.ver 0002827.000 0002c85.090 1000302.ver 10008e8.000 1
00020c4.090 00023ae.ver 00025e6.000 0002827.090 0002c85.ver 100030e.000 10008e8.090 1
00020c4.ver 00023af.000 00025e6.090 0002827.ver 0002c86.000 100030e.090 10008e8.ver 1
00020c5.000 00023af.090 00025e6.ver 0002828.000 0002c86.090 100030e.ver 10008fb.000 1
```

3. Create Test Folder

1. Create The test folder structure follows [Testing Standards for ucgmsim Git repositories](#)
2. Select 10 stations you want to test and cp corresponding waveforms files to the singel_files directory as below
3. Copy the source binary file 'BB_with_siteamp.bin' to the input folder
4. Run 'write_benchmark_csv(sample_bench_path)' function inside test_calculate_ims.py to generate 'new_im_sim_benchmark.csv', where 'sample_bench_path' is the folder we created in 1.4 Generate summary_benchmark: benchmark_im_sims. This function should only be run once for each binary file.

NOW you have all the input files ready

```
test
├── test_calculate_ims
│   ├── README
│   └── sample1
│       ├── input
│       │   ├── BB_with_siteamp.bin
│       │   ├── new_im_sim_benchmark.csv
│       │   └── single_files
│       │       ├── 00020d3.000
│       │       ├── 00020d3.090
│       │       ├── 00020d3.ver
│       │       ├── 2002199.000
│       │       ├── 2002199.090
│       │       ├── 2002199.ver
│       │       ├── CASH.000
│       │       ├── CASH.090
│       │       ├── CASH.ver
│       │       ├── CFW.000
│       │       ├── CFW.090
│       │       ├── CFW.ver
│       │       ├── DLX.000
│       │       ├── DLX.090
│       │       ├── DLX.ver
│       │       ├── EWZ.000
│       │       ├── EWZ.090
│       │       ├── EWZ.ver
│       │       ├── GRY.000
│       │       ├── GRY.090
│       │       ├── GRY.ver
│       │       ├── LSRC.000
│       │       ├── LSRC.090
│       │       ├── LSRC.ver
│       │       ├── PEAA.000
│       │       ├── PEAA.090
│       │       ├── PEAA.ver
│       │       ├── UNK.000
│       │       ├── UNK.090
│       │       └── UNK.ver
│       └── output
│           ├── ascii_darfield_im_sim
│           │   ├── ascii_darfield_im_sim.csv
│           │   ├── ascii_darfield_im_sim.info
│           │   └── stations
│           │       ├── 00020d3.csv
│           │       ├── 2002199.csv
│           │       ├── CASH.csv
│           │       ├── CFW.csv
│           │       ├── DLX.csv
│           │       ├── EWZ.csv
│           │       ├── GRY.csv
│           │       ├── LSRC.csv
│           │       ├── PEAA.csv
│           │       └── UNK.csv
│           └── binary_darfield_im_sim
│               ├── binary_darfield_im_sim.csv
│               ├── binary_darfield_im_sim.info
│               └── stations
│                   ├── 00020d3.csv
│                   ├── 2002199.csv
│                   ├── CASH.csv
│                   ├── CFW.csv
│                   └── DLX.csv
```



4. Run Pytest

Make sure you are currently under the test_calculate_ims folder, run:

```
$ pytest -v -s test_calculate_ims.py
```

```
yzh231@hypocentre ~/IM_calculation/test/test_calculate_ims (git)-[test] % pytest -v -s test_calculate_ims.py
===== test session starts =====
platform linux2 -- Python 2.7.14, pytest-3.2.2, py-1.4.34, pluggy-0.4.0 -- /usr/bin/python2.7
cachedir: ../../.cache
rootdir: /home/yzh231/IM_calculation, inifile:
plugins: cov-2.3.1
collecting 0 items
/home/yzh231/IM_calculation/calculate_ims.py
collected 6 items

test_calculate_ims.py::test_binary_script_calculate_ims python /home/yzh231/IM_calculation/calculate_ims.py /home/yzh231/IM_calcul
calculate_ims/sample1/output -i binary_darfield_im_sim -t s -n 2002199 GRY 00020d3 UNK CASH CFW DLX LSRC EWZ PEAA -p 0.01 0.2 0.5 1.
/home/yzh231/IM_calculation/test/test_calculate_ims/sample1/output/binary_darfield_im_sim/binary_darfield_im_sim.csv
Calculations are outputted to /home/yzh231/IM_calculation/test/test_calculate_ims/sample1/output/binary_darfield_im_sim

PASSED
test_calculate_ims.py::test_ascii_script_calculate_ims python /home/yzh231/IM_calculation/calculate_ims.py /home/yzh231/IM_calculat
ims/sample1/output -i ascii_darfield_im_sim -t s -n 2002199 GRY 00020d3 UNK CASH CFW DLX LSRC EWZ PEAA -p 0.01 0.2 0.5 1.0 3.0 4.0
```

CHECKPOINTING & SPLITTING A BIG SLURM

Responsible scripts

1. slurm header template: https://github.com/ucgmsim/slurm_gm_workflow/blob/master/templates/slurm_header.cfg
2. im_calc_slurm template: https://github.com/ucgmsim/slurm_gm_workflow/blob/master/templates/im_calc_sl.template
3. submit_hf.py that generates the slurm files: https://github.com/ucgmsim/slurm_gm_workflow/blob/master/scripts/submit_hf.py
4. checkpointing functions: https://github.com/ucgmsim/slurm_gm_workflow/blob/master/scripts/checkpoint.py

Checkpointing

Checkpointing is needed for IM_calculation due to large job size and limited running time on Kupe. Therefore, we implemented checkpointing to track the current progress of an im_calculation job, and carry on from where the job was interrupted by slurm.

Note, the checkpointing code relies on the input/output directory structure specified in the im_calc_al.template in the checkpoint branch. Failure to match the dir structure will result in runtime error. A quick fix would be modifying the template to suit your own dir structure.

Example:

(1) Simulation

Input/output structure defined in im_calc_al.template

```
echo __calculating simulations__
{% for sim_dir, sim_name, fault_name in sim_dirs %}
    time python2 $IMPATH/calculate_ims.py {{sim_dir}}/Acc/BB.bin b -o {{sim_dir}}/../../IM_calc/ -np 40 -i {{sim_name}} -r {{fault_name}} -t s -s
{% endfor %}
```

Actual input data structure:

```
melody.zhu@kupe01:/nesi/nobackup/nesi00213/RunFolder/Cybershake/v18p6_batched/v18p6_1k_under2p0G_ab/Runs> ls
Aratiatia ArielBank ArielEast Astrolabe01 Astrolabe02 Astrolabe03 Astrolabe05 Astrolabe07 Awakeri Barefell Bidwill Billys BlueLk BlueMtn BooBooEAST
```

The input binary file is under:

```
/nesi/nobackup/nesi00213/RunFolder/Cybershake/v18p6_batched/v18p6_1k_under2p0G_ab/Runs/BlueMtn/BB/Cant1D_v3-
midQ_OneRay_hfnp2mm+_rvf0p8_sd50_k0p045/BlueMtn_HYP28-31_S1514/Acc/BB.bin
```

The output **IM_calc** folder is under:

```
melody.zhu@kupe01:~/nesi/nobackup/nesi00213/RunFolder/Cybershake/v18p6_batched/v18p6_1k_under2p06_ab/Runs> ls ./BlueK/IM_calc
BlueK_HYP01-29_S1244 BlueK_HYP03-29_S1264 BlueK_HYP06-29_S1294 BlueK_HYP09-29_S1324 BlueK_HYP11-29_S1344 BlueK_HYP15-29_S1384 BlueK_HYP19-29_S1424 BlueK_HYP22-29_S1454 BlueK_HYP26-29_S1494 BlueK_HYP29-29_S1524
BlueK_HYP02-29_S1254 BlueK_HYP04-29_S1274 BlueK_HYP07-29_S1304 BlueK_HYP10-29_S1334 BlueK_HYP13-29_S1364 BlueK_HYP18-29_S1414 BlueK_HYP20-29_S1434 BlueK_HYP23-29_S1464 BlueK_HYP27-29_S1504
```

(2) Observed

Input/output structure defined in `im_calc_al.template`

```
echo calculating observed
{% for obs_dir, obs_name, fault_name in obs_dirs %}
time python2 $IMPATH/calculate_ims.py {{obs_dir}}/*/*accBB a -o {{obs_dir}}/../IM_calc/ -np {{np}} -i {{obs_name}} -r {{fault_name}} -t o -s
{% endfor %}
```

Actual input data structure:

```
melody.zhu@kupe01:~> ls test_obs/IMCalcExample/
2012p713691 2012p764736 2012p801609 2013p049577 2013p368016 2013p653606 2013p708602 2013p817946 2013p868761 2014p237547 2014p933966 2014p965622 2016p119534 2016p158394 2016p355841 2122842 3631755
```

The output **IM_calc** folder is under:

```
melody.zhu@kupe01:~> ls test_obs/IMCalcExample/IM_calc/
2012p713691 2012p764736 2012p801609 2013p049577 2013p368016 2013p653606 2013p708602 2013p817946 2013p868761
```

Splitting a big slurm

Splitting a big slurm script into several smaller slurms is needed due to the maximum number of lines allowed in a slurm script on Kupe.

Inside `submit_imcalc.py` The `-ml` argument specifies the maximum number of lines of python call to `calculate_ims.py`/`calculate_rrups.py`. **Header** and **footer** like `'#SBATCH --time=15:30:00', 'date'` etc are **NOT** included.

Say if the max number of lines allowed in a slurm script is 1000, and your (header + footer) is 30 lines, then the number `n` that you pass to `-ml` should be `0 < n <= 967`. eg. `-ml 967`.

Example:

We have 250 simulation dirs to run, by specifying `-ml 100` (100 python calls to `calculate_ims.py` per slurm script), we expect 3 sim slurm scripts to be outputted.(1-100, 100-200, 200-250)

We have 3 observed dirs to run, by specifying `-ml 100` (100 python calls to `calculate_ims.py` per slurm script), we expect 1 sim slurm scripts to be outputted.

We have 61 rrup files to run, by specifying `-ml 100` (100 python calls to `calculate_rrups.py` per slurm script), we expect 1 sim slurm scripts to be outputted.

Command to run checkpointing and splitting:

```
python submit_imcalc.py -obs ~/test_obs/IMCalcExample/ -sim runs/Runs -srf /nesi/nobackup/nesi00213/RunFolder/Cybershake/v18p6_batched/v18p6_exclude_1k_batch_6/Data/Sources -ll /scale_akl_nobackup/filesets/transit/nesi00213/StationInfo/non_uniform_whole_nz_with_real_stations-hh400_v18p6.ll -o ~/rrup_out -ml 1000 -e -s -i OtaraWest02_HYP01-21_S1244 Pahiatua_HYP01-26_S1244 -t 24:00:00
```

Output:

```
-rw-r--r-- 1 melody.zhu melody.zhu 48890 Jul 1 03:47 sim_im_calc_0.sl
-rw-r--r-- 1 melody.zhu melody.zhu 48674 Jul 1 03:47 sim_im_calc_100.sl
-rw-r--r-- 1 melody.zhu melody.zhu 31922 Jul 1 03:47 sim_im_calc_200.sl
-rw-r--r-- 1 melody.zhu melody.zhu 1329 Jul 1 03:47 obs_im_calc_0.sl
-rw-r--r-- 1 melody.zhu melody.zhu 23246 Jul 1 03:47 rrup_im_calc_0.sl
```

To submit the slurm script:

```
$cp test.sl /nesi/nobackup/nesi00213/tmp/auto_preproc
$sbatch test.sl
```

The reason that we have to run 'test.sl' under `'/nesi/nobackup/nesi00213/tmp/auto_preproc'` is otherwise slurm cannot find `machine.env` specified by the test.sl script:

```
source machine.env.sh
```

TODO

- Creation of semi-automatic slurm generation that will have all the calls to produce the results as needed.
- Progress printing statements
- Rrup calculation on a smaller station list - currently when generating the slurm script it does the full grid even for stations outside the domain

Notes

- Extensive re-writing of code needs to have smaller deliverables in the future, as this simplifies the integration.