# Documentation for Damage\_simulation.py

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## 1 Introduction

The damage function simulation is a key input into the pricing engine. Damages are represented in arrays of dimension nxp, where n = numstates and p = numperiods. The arrays are created by Monte Carlo simulation. Each array specifies for each state and time period a damage coefficient.

Up to a point, the Monte Carlo follows Pindyck (2012) 'Uncertain Outcomes and Climate Change Policy':

- 1. There is a gamma distribution for temperature
- 2. There is a gamma distribution for economic impact (conditional on temperature)

However, in addition, this program adds a probability of a tipping point (conditional on temperature). This probability is a decreasing function of the parameter 'peak<sub>t</sub>emp', conditional onatipping point

## 2 Damage Simulation Class

It is a class with a main function **simulate** which returns the simulated damage given a simulation method. The following methods are supported:

- 1. Pindyck displace gamma
- 2. Wagner-Weitzman normal
- 3. Roe-Baker
- 4. user-defined normal
- 5. user-defined gamma

## 2.1 Inputs and Outputs

### Inputs:

- tree : ('TreeModel' object) tree structure used
- ghg\_levels: (ndarray or list) end GHG level for each path
- **peak\_temp**: (float) tipping point parameter
- disaster\_tail : (float) curvature of tipping point
- tip\_on: (bool) flag that turns tipping points on or off
- $temp_map: (int)mappingfromGHGtotemperature$ 
  - 0: implies Pindyck displace gamma
  - 1: implies Wagner-Weitzman normal
  - 2: implies Roe-Baker
  - 3: implies user-defined normal
  - 4: implies user-defined gamma

**temp\_dist\_params**: (ndarray or list) if temp\_map is either 3 or 4, user needs to define the distribution parameters

maxh: (float) time parameter from Pindyck which indicates the time it takes for temp to get half way to its max value for a given level of ghg

cons\_growth: (float) yearly growth in consumption

#### Outputs:

The main output for this class is from function **simulate** which returns a 2-D array of damage indexed by x = number of final states and y = number of periods. Notice that a child only have one period and thus we can get a specific node given the final state and the period.

#### 2.2 Attributes

- tree: ('TreeModel' object) tree structure used
- ghg\_levels: (ndarray or list) end GHG level for each path
- **peak\_temp**: (float) tipping point parameter
- disaster\_tail: (float) curvature of tipping point
- tip\_on: (bool) flag that turns tipping points on or off
- $temp_map: (int)mappingfrom GHG to temperature$

- 0: implies Pindyck displace gamma
- 1: implies Wagner-Weitzman normal
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temp\_dist\_params: (ndarray or list) if temp\_map is either 3 or 4, user needs to define the distribution parameters

maxh: (float) time parameter from Pindyck which indicates the time it takes for temp to get half way to its max value for a given level of ghg

```
cons_growth : (float) yearly growth in consumptiond: (2-d array) simulated damage
```

#### 2.3 Methods

**\_gamma\_array**, **\_normal\_array**, **\_uniform\_array**: basic buildin functions to get random numbers of given dimension from gamma, normal and uniform distribution

```
def _gamma_array(self, shape, rate, dimension):
    return np.random.gamma(shape, 1.0/rate, dimension)

def _normal_array(self, mean, stdev, dimension):
    return np.random.normal(mean, stdev, dimension)

def _uniform_array(self, dimension):
    return np.random.random(dimension)
```

**\_sort\_array**: sort a given 2-D array to make the array is increasing in the periods. For example:

**\_normal\_simulation**: Draw random samples from normal distribution for mapping GHG to temperature given user defined parameter. **Inputs**:

- average: (ndarray or list): average temperature for each period
- std (ndarray or list): standard deviation for each period

#### Outputs:

• 1-D array of  $e^{simulated temperature}$ 

```
def _normal_simulation(self):
    """Draw random samples from normal distribution for mapping GHG to temperature
    user-defined distribution parameters.
    """
    assert self.temp_dist_params and len(self.temp_dist_params) == 2, "Normal distri
    ave, std = temp_dist_params
    n = len(ave)
    temperature = np.array([self._normal_array(ave[i],std[i], self.draws) for i in r
    return np.exp(temperature)
```

**\_gamma\_simulation**: Draw random samples from displaced gamma distribution for mapping GHG to temperature given user defined parameter.

Displaced gamma distribution is given by:

$$f(x; r, \lambda, \theta) = \frac{\lambda^r}{\Gamma(r)} (x - \theta)^{r-1} e^{-\lambda(x - \theta)}, x \ge \theta$$
 (1)

where  $\Gamma(r) = \int_0^{\inf} s^{r-1} e^{-s} ds$  is the gamma function. However, we used  $gamma(k, \theta) + displace$  to get the numerical result **Inputs**:

- k: (ndarray or list): shape parameter for each period
- theta (ndarray or list): scale parameter for each period
- displace (ndarray or list): displacement parameter for each period

#### **Outputs**:

• 1-D array of simulated temperature

\_pindyck\_simulation: Draw random samples for mapping GHG to temperature based on Pindyck. It is drawing from a gamma distribution but with the parameter given by pindyck

```
def _pindyck_simulation(self):
        """Draw random samples for mapping GHG to temperature based on Pindyck. The `p
        is the shape parameter from Pyndyck damage function, `pindyck_impact_theta` th
        from Pyndyck damage function, and `pindyck_impact_displace` the displacement p
        damage function.
        pindyck_temp_k = [2.81, 4.6134, 6.14]
        pindyck_temp_theta = [1.6667, 1.5974, 1.53139]
        pindyck_temp_displace = [-0.25, -0.5, -1.0]
        return np.array([self._gamma_array(pindyck_temp_k[i], pindyck_temp_theta[i], sel
                         + pindyck_temp_displace[i] for i in range(0, 3)])
_ww_simulation: Draw random samples for mapping GHG to temperature based on
Wagner-Weitzman. It is a drawing from a normal distribution with the parameters given by
Wagner-Weitzman.
    def _ww_simulation(self):
        """Draw random samples for mapping GHG to temperature based on Wagner-Weitzman
        ww_{temp_ave} = [0.573, 1.148, 1.563]
        ww_temp_stddev = [0.462, 0.441, 0.432]
        temperature = np.array([self._normal_array(ww_temp_ave[i], ww_temp_stddev[i], se
                                 for i in range(0, 3)])
        return np.exp(temperature)
_rb_simulation: It is drawing from a normal distribution with the parameters given by
Roe-Baker.
    def _rb_simulation(self):
        """Draw random samples for mapping GHG to temperature based on Roe-Baker."""
        rb_fbar = [0.75233, 0.844652, 0.858332]
        rb_sigf = [0.049921, 0.033055, 0.042408]
        rb_theta = [2.304627, 3.333599, 2.356967]
        temperature = np.array([self._normal_array(rb_fbar[i], rb_sigf[i], self.draws)
                         for i in range(0, 3)])
        return np.maximum(0.0, (1.0 / (1.0 - temperature)) - np.array(rb_theta)[:, np.ne
_pindyck_impact_simulation: It is drawing from a gamma distribution for the impact
with the parameter given by pindyck
   def _pindyck_impact_simulation(self):
        """Pindyck gamma distribution mapping temperature into damages."""
        # get the gamma in loss function
        pindyck_impact_k=4.5
        pindyck_impact_theta=21341.0
        pindyck_impact_displace=-0.0000746,
```

\_disaster\_simulation: Drawing random numbers from uniform distribution.

```
def _disaster_simulation(self):
    """Simulating disaster random variable, allowing for a tipping point to occur
    with a given probability, leading to a disaster and a `disaster_tail` impact o
    """
    disaster = self._uniform_array((self.draws, self.tree.num_periods))
    return disaster
```

\_disaster\_cons\_simulation: Generate TP\_damage in the paper from a gamma distribution with parameters  $\alpha = 1$  and  $\beta = dosaster\_tail$ .

```
def _disaster_cons_simulation(self):
    """Simulates consumption conditional on disaster, based on the parameter disas
    #get the tp_damage in the article which is drawed from a gamma distri with alp
    disaster_cons = self._gamma_array(1.0, self.disaster_tail, self.draws)
    return disaster_cons
```

<u>\_interpolation\_of\_temp</u>: for every temp in each period, modify it using a coefficient  $2 * (1 - 0.5^{timenow} timetoincrease half of the max regards to the current period.$ 

```
def _interpolation_of_temp(self, temperature):
     # for every temp in each period, modify it using a coff regards to the cur
return temperature[:, np.newaxis] * 2.0 * (1.0 - 0.5**(self.tree.decision_times[
```

**\_economic\_impact\_of\_temp**: calculate the economic impact of temperatures given temperature:

$$term_1 = \frac{-2*simulated\_impact*maxh*temp(foreachperiod)}{\log 0.5}$$
 
$$term_2 = con\_g - 2*simulated\_impact*temp*time_now$$
 
$$term_3 = \frac{2*gamma*maxh*temp*0.5(time_now/maxh)}{\log (0.5)}$$

and the final damage is  $e^{term_1+term_2+term_3}$ 

```
* temperature[:, np.newaxis] * 0.5**(self.tree.decision_times[1:] / self
return np.exp(term1 + term2 + term3)
```

\_tipping\_point\_update: Determine whether a tipping point has occurred, if so reduce consumption for all periods after this date. The step is as follows:

1. determine whether the tipping point is occurred by comparing the probability of survival and a random number generated from uniform distribution Where the probability of survival is:

$$prob_{survival} = \left[1 - \left(\frac{tmp}{tmp\_scale}\right)^{\frac{period\_len}{peak\_interval}}\right]$$

2. find unique final state and the periods that the diaster occurs and modify consumption after the point. (If a disaster happen more than once in a path, we only consider the influence of the first time.)

```
def _tipping_point_update(self, tmp, consump, peak_temp_interval=30.0):
    """Determine whether a tipping point has occurred, if so reduce consumption fo
    all periods after this date.
    11 11 11
   draws = tmp.shape[0]
   disaster = self._disaster_simulation()
   disaster_cons = self._disaster_cons_simulation()
   period_lengths = self.tree.decision_times[1:] - self.tree.decision_times[:-1]
   tmp_scale = np.maximum(self.peak_temp, tmp)
   ave_prob_of_survival = 1.0 - np.square(tmp / tmp_scale)
   prob_of_survival = ave_prob_of_survival**(period_lengths / peak_temp_interval) #
    # this part may be done better, this takes a long time to loop over
    # find unique row and the cols that the diaster occurs and modify consumption
   res = prob_of_survival < disaster
   rows, cols = np.nonzero(res)
   row, count = np.unique(rows, return_counts=True)
   first_occurance = zip(row, cols[np.insert(count.cumsum()[:-1],0,0)])
   for pos in first_occurance:
        consump[pos[0], pos[1]:] *= np.exp(-disaster_cons[pos[0]])
   return consump
```

\_run\_path: Calculate the distribution of damage for specific GHG-path. Varibles:

- tmp: smoothed temperature at a certain period
- consump: consumption at a certain period generated by \_economic\_impact\_of\_temp()
- **peak\_cons**: max consumption at a certain period generated by a constant growth rate: exp(constantgrowth\*timepassedfromthestartpoint)
- damage : 1.0 (consump / peak\_cons)
- weights: final\_states\_prob\*number\_of\_ draws

To determine what state does the damage be belong to, the code simply slice the damage array by the probability of a state occurs to classes. And then simply get the average within a class. **Output**:

• mean damage of the draws and return a 2-D array of damage

```
def _run_path(self, temperature):
        """Calculate the distribution of damage for specific GHG-path. Implementation
        the temperature and economic impacts from Pindyck [2012] page 6.
        11 11 11
        # Remark
        # -----
        # final states given periods can give us a specific state in that period since
        d = np.zeros((self.tree.num_final_states, self.tree.num_periods))
        tmp = self._interpolation_of_temp(temperature)
        consump = self._economic_impact_of_temp(temperature)
        peak_cons = np.exp(self.cons_growth*self.tree.decision_times[1:])
        # adding tipping points
        if self.tip_on:
            consump = self._tipping_point_update(tmp, consump)
        # sort based on outcome of simulation
        consump = self._sort_array(consump)
        damage = 1.0 - (consump / peak_cons)
        weights = self.tree.final_states_prob*(self.draws)
        weights = (weights.cumsum()).astype(int)
        d[0,] = damage[:weights[0], :].mean(axis=0)
        for n in range(1, self.tree.num_final_states):
            d[n,] = np.maximum(0.0, damage[weights[n-1]:weights[n], :].mean(axis=0))
        return d
simulate: main function of the class, multiprocessing run_path for a given method with
simulated temperature.
    def simulate(self, draws, write_to_file=True):
        """Create damage function values in 'p-period' version of the Summers - Zeckha
        Parameters
        _____
        draws : int
            number of samples drawn in Monte Carlo simulation.
        write_to_file : bool, optional
            wheter to save simulated values
```

```
Returns
_____
ndarray
    3D-array of simulated damages # it should be 2D : self.tree.num_final_stat
Raises
_____
ValueError
    If temp_map is not in the interval 0-4.
Note
Uses the :mod: `~multiprocessing` package.
11 11 11
dnum = len(self.ghg_levels)
self.draws = draws
self.peak_cons = np.exp(self.cons_growth*self.tree.decision_times[1:])
if self.temp_map == 0:
    temperature = self._pindyck_simulation()
elif self.temp_map == 1:
    temperature = self._ww_simulation()
elif self.temp_map == 2:
    temperature = self._rb_simulation()
elif self.temp_map == 3:
    temperature = self._normal_simulation()
elif self.temp_map == 4:
    temperature = self._gamma_simulation()
else:
    raise ValueError("temp_map not in interval 0-4")
pool = mp.Pool(processes=dnum)
self.d = np.array(pool.map(self._run_path, temperature))
if write_to_file:
    self._write_to_file()
return self.d
```