[Dacon] 블럭 장난감 제조 공정 최적화 경진대회

_(팀명)

2020년 월 일 (제출날짜)

- 1. 본 코드는 대회 참가를 돕고자 단순 예시를 작성한 것으로 참고용으로 사용바랍니다.
- 2. 본 코드는 자유롭게 수정하여 사용 할 수 있습니다.
- 3. 추가 모듈 보러가기: https://bit.ly/36MNs76 (https://bit.ly/36MNs76)

1. 라이브러리 및 데이터

Library & Data

In [1]:

```
import pandas as pd
import numpy as np
import multiprocessing
import warnings
from copy import deepcopy
from module.genome import Genome, genome_score
warnings.filterwarnings(action='ignore')
np.random.seed(777)
```

In [2]:

```
!python --version
print('Pandas : %s'%(pd.__version__))
print('Numpy : %s'%(np.__version__))
```

Pandas : 1.0.4 Numpy : 1.18.5

Python 3.6.10 :: Anaconda, Inc.

2. 데이터 전처리

Data Cleansing & Pre-Processing

In [3]:

입력하세요.

3. 탐색적 자료분석

Exploratory Data Analysis

In [4]:

입력하세요.

4. 변수 선택 및 모델 구축

Feature Engineering & Initial Modeling

In [5]:

```
CPU_CORE = multiprocessing.cpu_count() # 멀티프로세싱 CPU 사용 수
N_POPULATION = 100
                                     # 세대당 생성수
N_BEST = 10
                                     # 베스트 수
                                     # 자손 유전자 수
N_{CHILDREN} = 10
PROB MUTATION = 0.4
                                    # 돌연변이
REVERSE = False
                                     # 배열 순서 (False: ascending order, True: descending order)
                                    #초기점수
score_ini = 1e8
                                    # 입력 데이터 길이
input_length = 125
output_length_1 = 5
                                    # Event (CHECK_1~4, PROCESS)
                                    # MOL(0~5.5, step:0.5)
output_length_2 = 12
                                   # 히트레이어1 노드 수
h1 = 50
                                   # 히트레이어2 노드 수
h2 = 50
h3 = 50
                                   # 히트레이어3 노드 수
EPOCHS = 50
                                    # 반복 횟수
genomes = []
for _ in range(N_POPULATION):
   genome = Genome(score_ini, input_length, output_length_1, output_length_2, h1, h2, h3)
   genomes.append(genome)
try:
    for i in range(N_BEST):
       genomes[i] = best_genomes[i]
except:
   best_genomes = []
   for _ in range(5):
       genome = Genome(score_ini, input_length, output_length_1, output_length_2, h1, h2, h3)
       best_genomes.append(genome)
```

5. 모델 학습 및 검증

Model Tuning & Evaluation

- 1. PRT는 고정값 사용
- 2. Event A, Event B (MOL A, MOL B) 를 같은 값으로 제한
- 3. Event는 CHECK와 PROCESS 만 사용함
- 4. 목적 함수로 수요 부족분만 고려함
- 5. Event와 MOL에 대해 인공신경망 모델을 만들어 유전 알고리즘으로 학습

In [6]:

```
_gen = 1
core_history = []
igh_score_history = []
ean_score_history = []
hile n_gen <= EPOCHS:
      genomes = np.array(genomes)
      while len(genomes)%CPU_CORE != 0:
               genomes = np.append(genomes, Genome(score_ini, input_length, output_length_1, output_length_2
      genomes = genomes.reshape((len(genomes)//CPU_CORE, CPU_CORE))
      for idx, _genomes in enumerate(genomes):
               if __name__ == '__main__':
                        pool = multiprocessing.Pool(processes=CPU_CORE)
                        genomes[idx] = pool.map(genome_score, _genomes)
                        pool.close()
                        pool.join()
      genomes = list(genomes.reshape(genomes.shape[0]*genomes.shape[1]))
        # score에 따라 정렬
      genomes.sort(key=lambda x: x.score, reverse=REVERSE)
      # 평균 점수
      s = 0
      for i in range(N_BEST):
               s += genomes[i].score
      s /= N_BEST
      # Best Score
      bs = genomes[0].score
      # Best Model 추가
      if best_genomes is not None:
               genomes.extend(best_genomes)
      # score에 따라 정렬
      genomes.sort(key=lambda x: x.score, reverse=REVERSE)
      score_history.append([n_gen, genomes[0].score])
      high_score_history.append([n_gen, bs])
      mean_score_history.append([n_gen, s])
      # 결과 출력
      print('EPOCH #%s\thistory Best Score: \%s\thear Score: \%
      #모델 업데이트
      best_genomes = deepcopy(genomes[:N_BEST])
      # CHILDREN 생성
      for i in range(N_CHILDREN):
               new_genome = deepcopy(best_genomes[0])
               a_genome = np.random.choice(best_genomes)
               b_genome = np.random.choice(best_genomes)
               for j in range(input_length):
                        cut = np.random.randint(new_genome.w1.shape[1])
                        new_genome.w1[j, :cut] = a_genome.w1[j, :cut]
                        new_genome.w1[j, cut:] = b_genome.w1[j, cut:]
               for j in range(h1):
```

```
cut = np.random.randint(new_genome.w2.shape[1])
        new_genome.w2[j, :cut] = a_genome.w2[j, :cut]
        new_genome.w2[j, cut:] = b_genome.w2[j, cut:]
    for i in range(h2):
        cut = np.random.randint(new_genome.w3.shape[1])
        new_genome.w3[j, :cut] = a_genome.w3[j, :cut]
        new_genome.w3[j, cut:] = b_genome.w3[j, cut:]
    for j in range(h3):
        cut = np.random.randint(new_genome.w4.shape[1])
        new_genome.w4[j, :cut] = a_genome.w4[j, :cut]
        new_genome.w4[j, cut:] = b_genome.w4[j, cut:]
    for j in range(input_length):
        cut = np.random.randint(new_genome.w5.shape[1])
        new_genome.w5[j, :cut] = a_genome.w5[j, :cut]
        new_genome.w5[j, cut:] = b_genome.w5[j, cut:]
    for i in range(h1):
        cut = np.random.randint(new_genome.w6.shape[1])
        new_genome.w6[j, :cut] = a_genome.w6[j, :cut]
        new_genome.w6[j, cut:] = b_genome.w6[j, cut:]
    for i in range(h2):
        cut = np.random.randint(new_genome.w7.shape[1])
        new_genome.w7[j, :cut] = a_genome.w7[j, :cut]
        new_genome.w7[j, cut:] = b_genome.w7[j, cut:]
    for j in range(h3):
        cut = np.random.randint(new_genome.w8.shape[1])
        new_genome.w8[j, :cut] = a_genome.w8[j, :cut]
        new_genome.w8[i, cut:] = b_genome.w8[i, cut:]
   best_genomes.append(new_genome)
# 모델 초기화
genomes = []
for i in range(int(N_POPULATION / len(best_genomes))):
    for bg in best_genomes:
        new_genome = deepcopy(bg)
       mean = 0
        stddev = 0.2
        # 40% 확률로 모델 변형
        if np.random.uniform(0, 1) < PROB_MUTATION:</pre>
            new_genome.w1 += new_genome.w1 * np.random.normal(mean, stddev, size=(input_length, h
        if np.random.uniform(0, 1) < PROB_MUTATION:</pre>
            new_genome.w2 += new_genome.w2 * np.random.normal(mean, stddev, size=(h1, h2)) * np.r
        if np.random.uniform(0, 1) < PROB_MUTATION:
           new_genome.w3 += new_genome.w3 * np.random.normal(mean, stddev, size=(h2, h3)) * np.r
        if np.random.uniform(0, 1) < PROB_MUTATION:
           new_genome.w4 += new_genome.w4 * np.random.normal(mean, stddev, size=(h3, output_leng
        if np.random.uniform(0, 1) < PROB_MUTATION:
            new_genome.w5 += new_genome.w5 * np.random.normal(mean, stddev, size=(input_length, h
        if np.random.uniform(0, 1) < PROB_MUTATION:
           new_genome.w6 += new_genome.w6 * np.random.normal(mean, stddev, size=(h1, h2)) * np.r
        if np.random.uniform(0, 1) < PROB_MUTATION:
           new_genome.w7 += new_genome.w7 * np.random.normal(mean, stddev, size=(h2, h3)) * np.r
        if np.random.uniform(0, 1) < PROB_MUTATION:
            new_genome.w8 += new_genome.w8 * np.random.normal(mean, stddev, size=(h3, output_leng)
        genomes.append(new_genome)
```

```
if REVERSE:
    if bs < score_ini:
        genomes[len(genomes)//2:] = [Genome(score_ini, input_length, output_length_1, output_length]
else:
    if bs > score_ini:
        genomes[len(genomes)//2:] = [Genome(score_ini, input_length, output_length_1, output_length]
n_gen += 1
```

```
EPOCH #1
                History Best Score: 58350009.0 Best Score: 58350009.0
e: 73970253.2
               History Best Score: 52966224.0 Best Score: 52966224.0
EPOCH #2
e: 62719729.2
EPOCH #3
                History Best Score: 52966224.0 Best Score: 55631084.0
                                                                        Mean Scor
e: 61854906.8
EPOCH #4
                History Best Score: 52966224.0 Best Score: 56632663.0
                                                                        Mean Scor
e: 65462293.7
                History Best Score: 47669968.0 Best Score: 47669968.0
EPOCH #5
                                                                        Mean Scor
e: 63337793.2
EPOCH #6
                History Best Score: 46309053.0 Best Score: 46309053.0
                                                                        Mean Scor
e: 55622256.7
                History Best Score: 43432747.0 Best Score: 43432747.0
EPOCH #7
                                                                        Mean Scor
e: 52619787.3
                                                                        Mean Scor
EPOCH #8
                History Best Score: 43432747.0 Best Score: 43432747.0
e: 49620266.3
EPOCH #9
                History Best Score: 43189534.0 Best Score: 43189534.0
                                                                        Mean Scor
e: 46117971.3
EPOCH #10
                History Best Score: 43189534.0 Best Score: 43438367.0
                                                                        Mean Scor
e: 49911929.1
EPOCH #11
                History Best Score: 43189534.0 Best Score: 45688999.0
                                                                        Mean Scor
e: 51510627.3
EPOCH #12
                History Best Score: 43189534.0 Best Score: 45565105.0
e: 53140426.3
EPOCH #13
                History Best Score: 43189534.0 Best Score: 45282691.0
                                                                        Mean Scor
e: 51707521.9
EPOCH #14
                History Best Score: 43189534.0 Best Score: 43189534.0
                                                                        Mean Scor
e: 53988787.8
EPOCH #15
                History Best Score: 43189534.0 Best Score: 46317999.0
                                                                        Mean Scor
e: 49162608.5
                History Best Score: 43189534.0 Best Score: 46812804.0
EPOCH #16
                                                                        Mean Scor
e: 51259019.2
EPOCH #17
                History Best Score: 43189534.0 Best Score: 45962159.0
                                                                        Mean Scor
e: 49515542.9
EPOCH #18
                History Best Score: 43189534.0 Best Score: 43189534.0
                                                                        Mean Scor
e: 46884505.7
EPOCH #19
                History Best Score: 43189534.0 Best Score: 43189534.0
                                                                        Mean Scor
e: 43642192.3
EPOCH #20
                History Best Score: 43183585.0 Best Score: 43183585.0
                                                                        Mean Scor
e: 44836543.1
EPOCH #21
                History Best Score: 42403294.0 Best Score: 42403294.0
                                                                        Mean Scor
e: 43786573.3
EPOCH #22
                History Best Score: 42403294.0 Best Score: 43189534.0
e: 44314988.2
EPOCH #23
                History Best Score: 42403294.0 Best Score: 43024715.0
e: 45076389.4
EPOCH #24
                History Best Score: 42403294.0 Best Score: 44271847.0
e: 45533820.7
EPOCH #25
                History Best Score: 42403294.0 Best Score: 42414277.0
```

History Best Score: 42403294.0 Best Score: 43399136.0 Mean Scor

e: 43671763.8 EPOCH #26

e: 44210199.8

```
EPOCH #27
                                                        History Best Score: 42182941.0 Best Score: 42182941.0 Mean Scor
e: 43889956.3
EPOCH #28
                                                        History Best Score: 42182941.0 Best Score: 42182941.0 Mean Scor
e: 43636775.5
KeyboardInterrupt
                                                                                                                                                             Traceback (most recent call last)
<ipython-input-6-26c498a01ac4> in <module>
                                                        if __name__ == '__main__':
                                                                      pool = multiprocessing.Pool(processes=CPU_CORE)
                  13
     --> 14
                                                                                 genomes[idx] = pool.map(genome_score, _genomes)
                  15
                                                                       pool.close()
                  16
                                                                      pool.ioin()
~\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda3\manaconda
nc, iterable, chunksize)
               264
                                                         in a list that is returned.
              265
 --> 266
                                                                return self._map_async(func, iterable, mapstar, chunksize).get()
              267
              268
                                          def starmap(self, func, iterable, chunksize=None):
~\makepanaconda3\makepanvs\makepattensorflow2.0\makepallib\multiprocessing\makeppool.py in get(self, ti
meout)
              636
              637
                                          def get(self. timeout=None):
                                                                self.wait(timeout)
 --> 638
              639
                                                         if not self.ready():
                                                                      raise TimeoutError
              640
~\maconda3\makenvs\makentensorflow2.0\makeplib\makepmultiprocessing\makeppool.py in wait(self, ti
meout)
              633
              634
                                          def wait(self, timeout=None):
 --> 635
                                                                self._event.wait(timeout)
              636
                                          def get(self, timeout=None):
              637
~\makepanaconda3\makepanvs\makepattensorflow2.0\makepattenlib\makepatthreading.py in wait(self, timeout)
              549
                                                                      signaled = self._flag
              550
                                                                       if not signaled:
--> 551
                                                                                                 signaled = self._cond.wait(timeout)
              552
                                                                       return signaled
              553
~\makepanaconda3\makepanvs\makepattensorflow2.0\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\makepattentlib\mak
              293
                                                                                       # restore state no matter what (e.g., KeyboardInterr
upt)
              294
                                                                       if timeout is None:
 --> 295
                                                                                                 waiter.acquire()
              296
                                                                                     gotit = True
              297
                                                                       else:
KeyboardInterrupt:
```

6. 결과 및 결언

Conclusion & Discussion

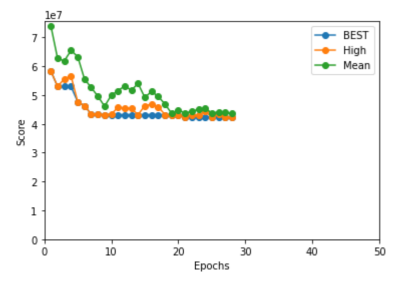
결과 그래프

In [7]:

```
import matplotlib.pyplot as plt

# Score Graph
score_history = np.array(score_history)
high_score_history = np.array(high_score_history)
mean_score_history = np.array(mean_score_history)

plt.plot(score_history[:,0], score_history[:,1], '-o', label='BEST')
plt.plot(high_score_history[:,0], high_score_history[:,1], '-o', label='High')
plt.plot(mean_score_history[:,0], mean_score_history[:,1], '-o', label='Mean')
plt.legend()
plt.xlim(0, EPOCHS)
plt.ylim(bottom=0)
plt.xlabel('Epochs')
plt.ylabel('Score')
plt.show()
```



Submission 파일 만들기

In [8]:

```
# 재立 계산
from module.simulator import Simulator
simulator = Simulator()
order = pd.read_csv('module/order.csv')
submission = best_genomes[0].predict(order)
_, df_stock = simulator.get_score(submission)

# PRT 개수 계산
PRTs = df_stock[['PRT_1', 'PRT_2', 'PRT_3', 'PRT_4']].values
PRTs = (PRTs[:-1] - PRTs[1:])[24*23:]
PRTs = np.ceil(PRTs * 1.1)
PAD = np.zeros((24*23+1, 4))
PRTs = np.append(PRTs, PAD, axis=0).astype(int)

# Submission 파일에 PRT 일록
submission.loc[:, 'PRT_1':'PRT_4'] = PRTs
submission.to_csv('Dacon_baseline.csv', index=False)
```

점수 향상 팁

해당 코드는 단순한 모델로 다음 방법으로 점수 향상을 꾀할 수 있습니다.

- 1. 성형 공정 2개 라인을 따로 모델링
- 2. CHANGE, STOP 이벤트 활용
- 3. 수요 초과분 외 다양한 양상을 반영하는 목적함수
- 4. 유전 알고리즘 외 효율적인 학습 기법

In []: