# 大数据导论期末 Project

组员:

- 赵博弘 11612706
- 郭舒雨 11712932

老师,很抱歉,这个PDF是之后才调好的,不知道可不可以不算迟交

# 背景信息

2020年,1月下旬,新冠肺炎开始在亚洲地区产生,该肺炎具有致死率低,传播快的特点,如不进行即时的防治措施,感染人数将随着时间的推移快速增长并呈现爆发之势。而韩国由于没有及时地进行相应的隔离措施,导致其成为亚洲的重灾区。韩国的总人口约为五千一百万,在四月累计感染人数达到了一万以上,累计死亡人数达到了两百人次。在本次project中,我们收集到了韩国在疫情爆发期间(1月20日至4月6日)的信息,包括病人信息,天气,城市教育水平以及医疗负荷水平,对韩国的总体疫情发展以及病人个体的康复情况都进行了数据分析,试图找出一些规律。

# Task1:数据预处理

像这类政府收集的信息,一般来说都会以表格的方式存储,数据结构比较简陋,且不说数据缺失是家常便饭,甚至可能混进一些不合法的数据。比如数值型数据中出现其他符号,单词拼写错误,出现与列属性表达规范不同的数据等。

# 预处理之前

- 检查非法数据
- 1. 对于数值型的,导入过程中以数值类型导入,查看是否会出现报错
- 2. 对于字符串类型的,在本项目中一般情况为分类数据,直接使用 DataFrame 的 value\_counts 方法,基本上每个 label 都会有比较多的数量,那么只需要检查数量为 1 的数据是否非法即可。

检查到的非法数据,我们会根据情况判断是否丢弃,或者直接在元数据表中修改。

# 缺失值填补

- Infection case: 使用 etc 表示未知数结
- contact number: 使用 0 填补
- infected by: 使用-1表示未知的传染者
- age, birth\_year: 这两个属性有关联性,所以一起填补。观察数据得出,有一部分人没有 birth\_year,而且仅在那一部分人中,存在二值都缺失的人。于是首先对于两者都缺失的人的 birth\_year,以该人所在 province 的 mean 进行填补,然后可以推导得出这一部分人的 age, 此时只剩下一部分人有 age 且没有 birth\_year。最后以 age 分组,这些人的 birth year 以此人 age 组内的 birth year 的 mean 进行填补。
- country: 直接丢掉这个列
- city: 因为每个人的 province 是完整的,默认认为这个 info 表中的地点为该人的户籍地点,所以直接使用每个人的 route 中去过最多的城市来填补,对于 route 中没有出现的人,直接使用该 province 中最多的 city 来填补。
- desease: 进行了转换,有病用1代表,其他用0代表。
- 日期转换:将表中字符串类型的日期转换成内置的日期类

# 新生成数据

• start\_date: 我们想如果能知道一个人的具体的患病日期(而不仅仅是 symptom\_onset\_date 和 confirmed\_date ) 是极为关键的。这样就可以推导出比如像该患者等了多久才去就医的这个时长,过程中有没有去过其他地方进行传播等,对后期的 回归预测会有很大帮助。

首先看该人有没有 symptom\_onset\_date ,如果没有就选择 confirmed\_date ,作为 0day 。然后看在 0day 之前该人的 route 中有没有去过其他地方,对应的会有一系列的时间节点,如果有就从其中选择时间节点与 0day 最接近的那个日期作为 start\_date 。如果没有,就选择 0day 作为 start\_date 。

• ..

```
"TOAU_EAL AUCUTETOAU
%autoreload 2
import sys
import random
import folium
from folium import plugins
import pandas as pd
import numpy as np
import matplotlib.pyplot as plt
import seaborn as sns
import preproc as pp
from sklearn.cluster import KMeans
from sklearn.ensemble import RandomForestClassifier
from sklearn.metrics import f1 score as f1
from sklearn.metrics import confusion matrix as con
from sklearn.naive_bayes import MultinomialNB
from sklearn.model_selection import train_test_split
from sklearn.linear_model import LinearRegression
from sklearn import metrics
class COVID(object):
    def init (self):
        self.DATA PATH = "../coronavirusdataset/"
        self.patient route = pp.patient route(self.DATA PATH)
        self.case, self.group_map = pp.case(self.DATA_PATH)
        self.patient_info = pp.patient_info(self.DATA PATH)
        self.patient num = self.patient info.shape[0]
        self.time = pp.time(self.DATA PATH)
        self.region = pp.region(self.DATA_PATH)
        self.pre_proc()
    def pre_proc(self):
        # >>> patient info:
        # fill missing value: infection case, use etc
        self.patient info["infection case"].fillna("etc", inplace=True)
        # fill missing value: contact_number, use 0
        self.patient_info["contact_number"].fillna(0, inplace=True)
         ^{2} fill missing value: infected by, use -1
        self.patient_info["infected_by"].fillna(-1, inplace=True)
        # fill missing value: sex, use ratio of all, generate a shuffle list then fill in the
blank
        sex_count = dict(self.patient_info["sex"].value_counts())
        sex num = sex count["female"] + sex count["male"]
        un_sex_male = int((self.patient_num - sex_num) * sex_count["male"] / sex_num)
        un_sex_female = int((self.patient_num - sex_num) - un_sex_male)
        fill_sex_list = ["male" for _ in range(0, un_sex_male)] + ["female" for _ in range(0, un_sex
_female)]
        random.shuffle(fill sex list)
        self.patient_info.loc[self.patient_info["sex"].isnull(), "sex"] = fill_sex_list
        # fill missing value: age, use mean of province
        # # 1. fill birth year
        all_age = self.patient_info.birth_year.mean()
        province_age = self.patient_info.groupby('province').birth_year.mean()
        province age.loc[province age.isnull()] = all age
        fill_birth_list_idx = self.patient_info.loc[(self.patient_info["birth_year"].isnull()) & (s
elf.patient info["age"].isnull()), "birth year"].index
        fill birth list = list(province age[self.patient info.loc[fill birth list idx, "province"]]
        self.patient_info.loc[(self.patient_info["birth_year"].isnull()) & (self.patient_info["age"
].isnull()), "birth_year"] = fill_birth_list
        # # 2. fill age
        fill age list idx = self.patient info.loc[self.patient info["age"].isnull(), "age"].index
        cal_year_list = np.array(list(self.patient_info.loc[fill_age_list_idx, "birth_year"]))
        cal_year_list = (2020 - cal_year_list) / 10
        cal_year_list = cal_year_list.astype(int) * 10
cal_year_list = [str(cal_year_list[i]) + 's' for i in range(cal_year_list.shape[0])]
        self.patient_info.loc[self.patient_info["age"].isnull(), "age"] = cal_year_list
        # fill birth again
        birth mean = self.patient info.groupby('age').birth year.mean()
        fill_birth_list_idx = self.patient_info.loc[self.patient_info["birth_year"].isnull(), "birt
h year"].index
        fill_birth_list = list(birth_mean[self.patient_info.loc[fill_birth_list_idx, "age"]])
        self.patient info.loc[self.patient info["birth year"].isnull(), "birth year"] = fill birth
list
        # drop country
        solf nationt info dron/solumns=["sountry"] inplace=mmuo)
```

```
serr.patrent_into.drop(corumns-[ country ], inprace-True)
        # fill missing value: city
        self.patient_info.loc[self.patient_info["city"] == "etc", "city"] = None
        self.patient_route["most_city"] = self.patient_route.groupby('patient_id').city.transform(
            lambda x: x.value_counts().index[0])
        fill_city_list_idx = self.patient_info.loc[self.patient_info["city"].isnull(), "city"].inde
Х
        route_most_city = self.patient_route.groupby('patient_id').most_city.max()
        fill_city_list_pid = self.patient_info.loc[fill_city_list_idx, "patient_id"]
        for idx, pid in fill_city_list_pid.iteritems():
            if pid in route most city.index:
                self.patient_info.loc[idx, "city"] = route_most_city[pid]
        self.patient info["tmp"] = self.patient info.groupby('province').city.transform(
            lambda x: x.value counts().index[0])
        fill_city_list_idx = self.patient_info.loc[self.patient_info["city"].isnull(), "city"].inde
        prov_most_city = self.patient_info.groupby('province').tmp.max()
        fill city_list_pid = self.patient_info.loc[fill_city_list_idx, "patient_id"]
        for idx, pid in fill_city_list_pid.iteritems():
            # print(self.patient_info.loc[idx, "province"])
            self.patient_info.loc[idx, "city"] = prov_most_city[self.patient_info.loc[idx,
"province"]]
        # fill last city and province
        ttp = self.patient route.groupby("patient id").last().reset index()
        last_index = self.patient_info.loc[self.patient_info["patient_id"].isin(ttp["patient_id"])]
.index
        last index = np.array(last index)
        for idx in range(len(last_index)):
            self.patient info.loc[self.patient info.patient id == ttp.loc[idx, "patient id"], "last
_city"] = ttp.loc[idx, "city"]
           self.patient info.loc[self.patient info.patient id == ttp.loc[idx, "patient id"], "last
_province"] = ttp.loc[idx, "province"]
        null_last_idx = self.patient_info.loc[self.patient_info["last_city"].isnull()].index
        self.patient_info.loc[null_last_idx, "last_city"] = self.patient_info.loc[null_last_idx, "c
ity"]
        self.patient_info.loc[null_last_idx, "last_province"] = self.patient_info.loc[null_last_idx
, "province"]
        # fill disease
        self.patient_info["disease_b"] = 1
        self.patient_info.loc[self.patient_info["disease"].isnull(), "disease_b"] = 0
        self.patient_info["disease_b"].value_counts()
        # date formatting
        self.patient info.loc[776, "symptom onset date"] = None
        self.patient info["confirmed date"] = pd.to datetime(self.patient info.confirmed date)
        self.patient_info["symptom_onset_date"] = pd.to_datetime(self.patient_info.symptom_onset_da
te)
        self.patient info["released date"] = pd.to datetime(self.patient info.released date)
        self.patient info["deceased_date"] = pd.to_datetime(self.patient_info.deceased_date)
        self.time["date"] = pd.to_datetime(self.time.date)
        # >>> patient_route:
        self.patient_route["date"] = pd.to_datetime(self.patient_route.date)
        self.patient_info["start_date"] = pd.NaT
        for i, row in self.patient info.iterrows():
            pid = row["patient id"]
            dating = self.patient_route[self.patient_route["patient_id"] == pid]
            self.patient info.loc[self.patient info["patient id"] == pid, "group"] = self.group map
[row["infection_case"]]
            if row["symptom_onset_date"] is not pd.NaT:
                if dating.shape[0] == 0:
                    self.patient_info.loc[i, "start_date"] = row["symptom_onset_date"]
                    continue
                dating_b = dating[dating["date"] <= row["symptom_onset_date"]]</pre>
                if dating_b.shape[0] == 0:
                    self.patient info.loc[i, "start date"] = row["symptom onset date"]
                self.patient_info.loc[i, "start_date"] = dating_b.tail(1).reset_index().at[0, "date
            else:
                if dating.shape[0] == 0:
                    self.patient_info.loc[i, "start_date"] = row["confirmed_date"]
                    continue
                dating b = dating[dating["date"] <= row["confirmed date"]]</pre>
                if dating_b.shape[0] == 0:
                    self.patient_info.loc[i, "start_date"] = row["confirmed_date"]
                self.patient_info.loc[i, "start_date"] = dating_b.tail(1).reset_index().at[0, "date
```

```
print(self.patient_info.isnull().sum())
    # fill missing value: sym date
    # self.patient_info.to_excel("./tmp/patient_info.xlsx")

covid_model = COVID()

patient_id 0
```

0 sex birth\_year 0 0 age province 0 city infection case 0 infected\_by 0  ${\tt contact\_number}$ 0 disease 3110 symptom\_onset\_date 2683 confirmed\_date 0 released\_date 2147 deceased\_date 3072 0 state group 0 tmp 0 last\_city last province 0 disease b 0 start date dtype: int64

# 统计信息与可视化展示

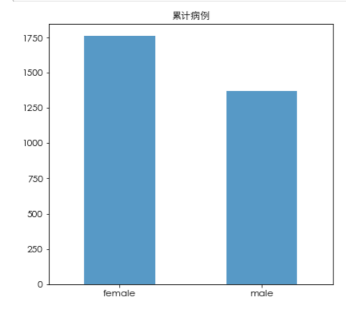
# 性别分布

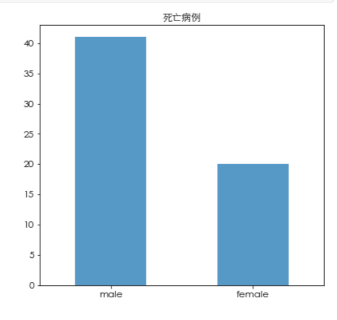
可见女性患者略多于男性患者,男性的死亡人数多于女性。

## In [35]:

```
# plot: sex
plt.rcParams['font.family']=['Heiti TC']

deceased = covid_model.patient_info[covid_model.patient_info["state"] == "deceased"]
plt.figure()
plt.rcParams["figure.figsize"] = (14.0, 6.0)
plt.subplot(121)
plt.title("累计病例")
covid_model.patient_info["sex"].value_counts().plot(kind="bar", alpha=0.75, rot=0, fontsize=12)
plt.subplot(122)
plt.title("死亡病例")
deceased["sex"].value_counts().plot(kind="bar", alpha=0.75, rot=0, fontsize=12)
plt.show()
```



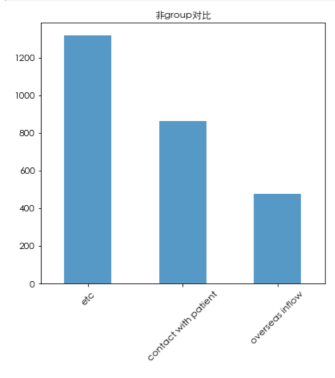


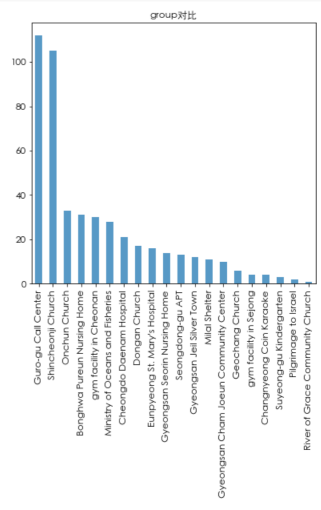
# 案例分布

- etc 这种未知原因还是占很大一部分比例的。所以与 contact with patient 和 overseas inflow 这两个基数也很大的案例放在一起对比。
- 而对于其他的案例,通常是群体集会行为/前往人群密集区,大部分是在医院接触了其他患者以及教堂的公众集会。

#### In [68]:

```
# other infection case
plt.figure()
plt.rcParams["figure.figsize"] = (14.0, 6.0)
plt.subplot(121)
plt.title("非group对比")
covid_model.patient_info["infection_case"].value_counts()[:3].plot(kind="bar", alpha=0.75, rot=45, fontsize=12)
plt.subplot(122)
plt.title("group对比")
covid_model.patient_info["infection_case"].value_counts()[3:].plot(kind="bar", alpha=0.75, rot=90, fontsize=12)
plt.show()
```

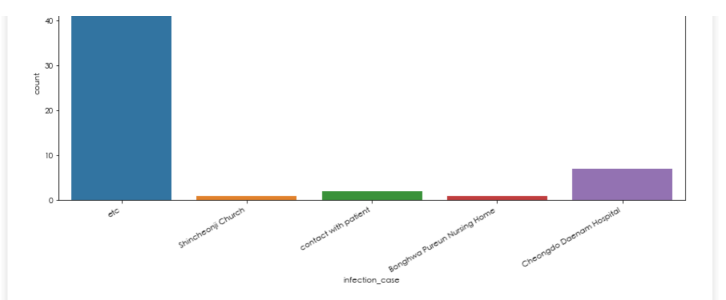




• 死亡者中感染状况为非群体感染的人数较远多于其他情况.

#### In [53]:

```
deceased_case=sns.countplot(x="infection_case", data=deceased)
plt.gcf().autofmt_xdate()
```



# 城市分布

以下二图为病患的城市分布和热力图

#### In [162]:

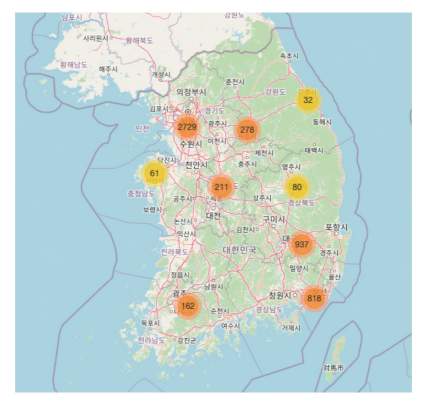
```
# define the kr map
loc = [36.4268093, 126.5512681]
kr_map = folium.Map(location=loc, zoom_start=7)

# add all patient
incidents = plugins.MarkerCluster().add_to(kr_map)
for lat, lng, label, in zip(covid_model.patient_route.latitude, covid_model.patient_route.longitude
, covid_model.patient_route.province):
    folium.Marker(
        location=[lat, lng],
        icon=None,
        popup=label,
        ).add_to(incidents)

# display kr map
kr_map
```

#### Out[162]:

Make this Notebook Trusted to load map: File -> Trust Notebook



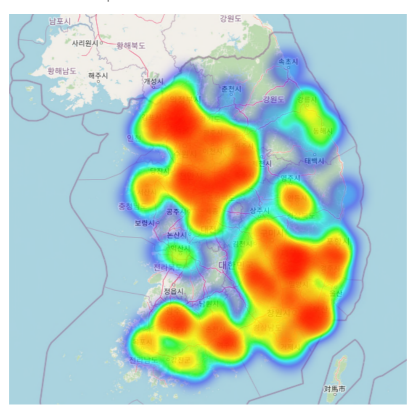
# In [110]:

```
# define the kr map
loc = [36.4268093, 126.5512681]
kr_map = folium.Map(location=loc, zoom_start=7)

# add heatmap
heatdata = covid_model.patient_route[['latitude','longitude']].values.tolist()
plugins.HeatMap(heatdata).add_to(kr_map)
kr_map
```

#### Out[110]:

Make this Notebook Trusted to load map: File -> Trust Notebook



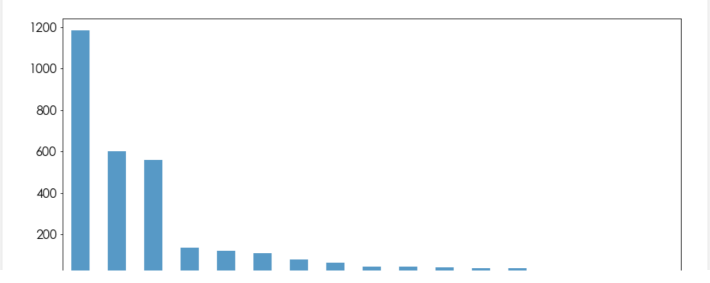
# 病患城市分布

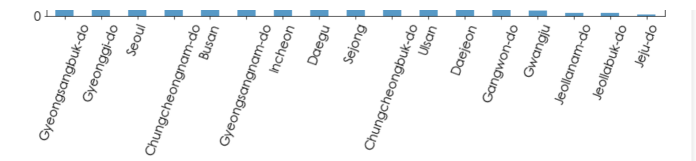
#### In [60]:

```
covid_model.patient_info["province"].value_counts().plot(kind="bar", alpha=0.75, rot=70, fontsize=1
6)
```

## Out[60]:

<matplotlib.axes.\_subplots.AxesSubplot at 0x7f8ee78ab090>

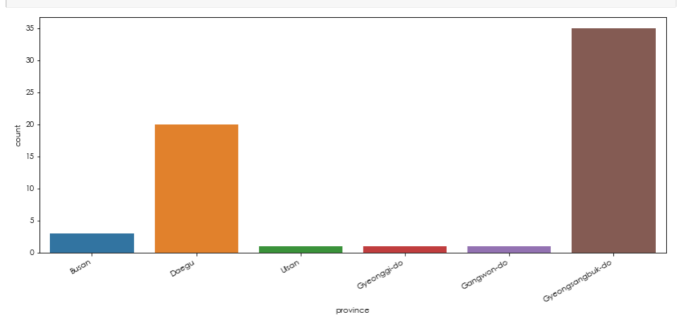




死亡的病人中,最多的是来自Gyeongsangbuk-do和Daegu这两个地方

#### In [61]:

```
deceased_province=sns.countplot(x="province", data=deceased)
plt.gcf().autofmt_xdate()
```



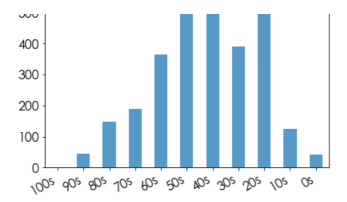
# 年龄分布

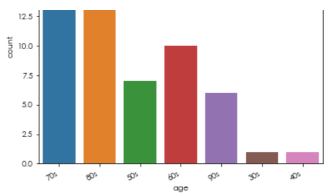
- 病患主要集中于中老年人和20多岁的年轻人。
- 死亡的病人多为60,70,80代,因此我们可以粗略看出,此病毒对于免疫力低下的老龄人口具有较大的威胁性。死亡的病人多为60,70,80代,因此我们可以粗略看出,此病毒对于免疫力低下的老龄人口具有较大的威胁性。

#### In [89]:

```
count = covid_model.patient_info["age"].value_counts()
bar = ['100s','90s','80s','70s','60s','50s','40s','30s','20s','10s','0s']
plt.figure()
plt.rcParams["figure.figsize"] = (14.0, 6.0)
plt.subplot(121)
plt.title("累计病例")
count[bar].plot(kind="bar", alpha=0.75, rot=0, fontsize=16)
plt.subplot(122)
plt.title("死亡病例")
deceased_age=sns.countplot(x="age", data=deceased)
plt.gcf().autofmt_xdate()
plt.show()
```







# 时间分布的数据

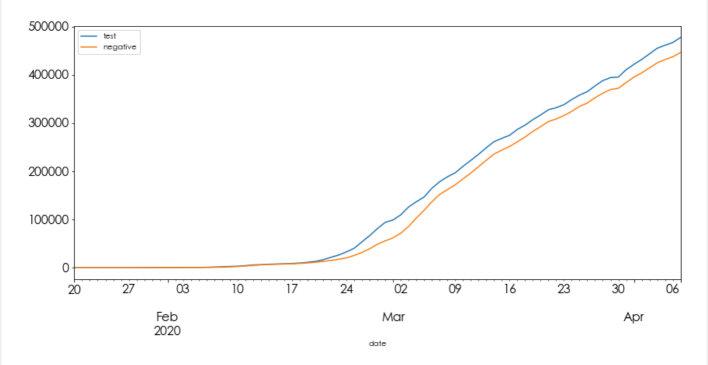
#### 病毒检测

可见韩国从三月开始对病毒的检测才踏入正轨。

# In [119]:

```
timep = covid_model.time.set_index("date")
plt.figure()
plt.rcParams["figure.figsize"] = (14.0, 6.0)
timep[["test", "negative"]].plot(fontsize=16)
plt.show()
```

<Figure size 1008x432 with 0 Axes>

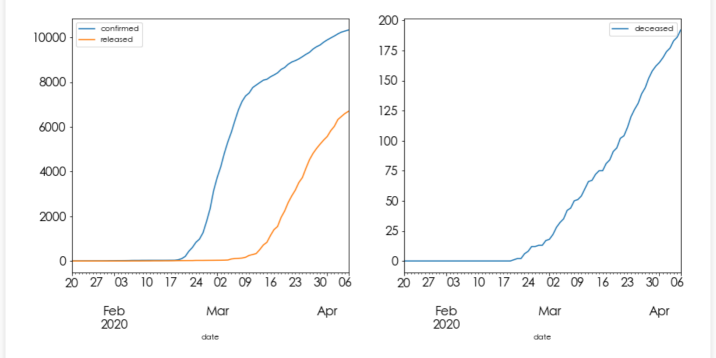


根据累计死亡人数和累计确诊人数的时间序列图,我们可以看出韩国感染人数从2月下旬开始呈现爆发性的增长,到3月中旬增长速度较为缓和,而死亡人数增速一直呈现上升趋势。根据累计死亡人数和累计确诊人数的时间序列图,我们可以看出韩国感染人数从2月下旬开始呈现爆发性的增长,到3月中旬增长速度较为缓和,而死亡人数增速一直呈现上升趋势。

#### In [121]:

```
plt.figure()
figure, axes = plt.subplots(1, 2)
plt.rcParams["figure.figsize"] = (14.0, 6.0)
timep[["confirmed", "released"]].plot(fontsize=16, ax=axes[0])
timep[["deceased"]].plot(fontsize=16, ax=axes[1])
plt.show()
```

<Figure size 1008x432 with 0 Axes>



# 特征工程

接下来,我们根据patientInfo中的病人籍贯地点对该地点教育水平进行合并,根据PatientRoute的中病人接受治疗的地点对该地点医疗负荷数据进行了合并。我们提取出了状态为released的病人数据,将出院时间和确诊时间进行相减粗略得到病人康复所需时间,并结合并性别、年龄、群体感染与否、学术水平以及医疗负荷水平进行了相关系数的计算。我们发现病人康复所需时间与其他变量之间的相关性非常弱,因此我们认为这些变量对于病人康复所需时间不具有明显的解释性。

#### In [143]:

```
region = covid model.region.copy()
info = covid_model.patient_info.copy()
time = covid_model.time.copy()
info.loc[37, "city"] = "Songpa-gu"
info.loc[106, "city"]="Geumcheon-gu"
info.loc[269, "city"]="Gangnam-gu"
info.loc[385, "city"]="Guro-gu"
info.loc[562, "province"]="Chungcheongnam-do"
info.loc[771, "city"]="Seo-gu"
info.loc[1812, "city"]="Mokpo-si"
info.loc[562, "last_province"]="Chungcheongnam-do"
region["last province"]=region["province"]
region["last_city"]=region["city"]
new sex=pd.get dummies(info["sex"])
info["sex"]=new_sex["male"]#### male 为1
a = np.array(info["group"])
group=a.astype(int)
info["group"]=group#### group 为1
for i in range(len(info["age"])):
    info.loc[i, "age"] = int(info["age"][i].strip("s"))
timeage=pd.read_csv("../coronavirusdataset/TimeAge.csv")
timegender=pd.read csv("../coronavirusdataset/TimeGender.csv")
timeprovince=pd.read_csv("../coronavirusdataset/TimeProvince.csv")
part1=region[["province","city","elementary_school_count","kindergarten_count","university count",
"academy_ratio"]]
part2=region[["last_province","last_city","elderly_population_ratio","elderly_alone_ratio","nursing
_home_count"]]
infol=pd.merge(info,part1,on = ["province","city"],how = "left")
final_info=pd.merge(info1,part2,on = ['last_province','last_city'],how = 'left')
released=final_info[final_info["released_date"].isnull()!=True]
released=released[released["state"]=="released"]
rctime=released["released_date"]-released["confirmed_date"]
rctime=rctime.dt.days[0:len(rctime)]
```

#### Out[143]:

|                          | sex      | group    | elementary_school_count | kindergarten_count | university_count | academy_ratio | elderly_po <sub>l</sub> |
|--------------------------|----------|----------|-------------------------|--------------------|------------------|---------------|-------------------------|
| sex                      | 1.000000 | 0.050915 | 0.009009                | 0.009470           | -0.180372        | 0.076846      |                         |
| group                    | 0.050915 | 1.000000 | 0.063677                | -0.018200          | -0.227172        | 0.083858      |                         |
| elementary_school_count  | 0.009009 | 0.063677 | 1.000000                | 0.942532           | -0.005065        | 0.355380      |                         |
| kindergarten_count       | 0.009470 | 0.018200 | 0.942532                | 1.000000           | 0.149475         | 0.308139      |                         |
| university_count         | 0.180372 | 0.227172 | -0.005065               | 0.149475           | 1.000000         | -0.256601     |                         |
| academy_ratio            | 0.076846 | 0.083858 | 0.355380                | 0.308139           | -0.256601        | 1.000000      |                         |
| elderly_population_ratio | 0.010878 | 0.002591 | -0.531670               | -0.575729          | -0.114868        | -0.464459     |                         |
| elderly_alone_ratio      | 0.002265 | 0.026354 | -0.442323               | -0.496959          | -0.199616        | -0.431836     |                         |
| nursing_home_count       | 0.065492 | 0.003719 | 0.653121                | 0.604776           | -0.154123        | 0.521780      |                         |
| recoverytime             | 0.019096 | 0.021578 | -0.029925               | -0.004614          | 0.081694         | -0.080391     |                         |
|                          |          |          |                         |                    |                  |               |                         |

我们接下来对病人是否死亡或者出院进行了分类预测。主要利用性别、年龄、群体感染与否、学术水平以及医疗负荷水平等变量。

### In [144]:

deceased 0.952646
Name: state, dtype: float64
released 0.947305
deceased 0.052695
Name: state, dtype: float64

# 朴素贝叶斯模型

### In [145]:

```
classifier = MultinomialNB().fit(x_train,y_train)
test_pred = classifier.predict(x_test)
train_pred = classifier.predict(x_train)
```

#### 模型性能评估

根据朴素贝叶斯模型的分类结果,可以看出运用该种方法分类的precision较低,但是recall召回率较高,但是有较多实际为released的 病人被误分为deceased。若按照这种方法为院方进行分类估计,易给医院造成较大的负担和压力。

```
In [148]:
```

```
print(metrics.confusion_matrix(y_test, test_pred), "\n")
print(metrics.classification_report(y_test,test_pred))
[[ 17 0]
 [ 96 246]]
             precision recall f1-score support
                  0.15
                           1.00
                                     0.26
                                                 17
   deceased
   released
                  1.00
                           0.72
                                     0.84
                                                342
   accuracy
                                     0.73
                                                359
                            0.86
                  0.58
                                     0.55
                                                359
  macro avg
weighted avg
                  0.96
                           0.73
                                     0.81
                                                359
```

# 随机森林

#### In [151]:

0.9749303621169917

```
rf = RandomForestClassifier(class_weight='balanced',random_state=1)
###返回由训练集训练成的模型对验证集预测的结果
test_pred = rf.fit(x_train,y_train).predict(x_test)
###打印混淆矩阵
print('\n'+'混淆矩阵: ')
print(con(y_test,test_pred))
###打印测试准确率
print('\n'+'Accuracy:')
print(rf.score(x_test,y_test), "\n")
print(metrics.classification_report(y_test,test_pred))

混淆矩阵:
[[ 13     4]
    [ 5     337]]
Accuracy:
```

#### precision recall f1-score support deceased 0.72 0.76 0.74 17 released 0.99 0.99 0.99 342 accuracy 0.97 359 0.86 0.88 0.86 359 macro avg 0.98 0.97 0.98 359 weighted avg

对比朴素贝叶斯,我们可以看到运用了随机森林的算法分类的效果更好。更多的实际出院病人被进行了正确的预测分类。

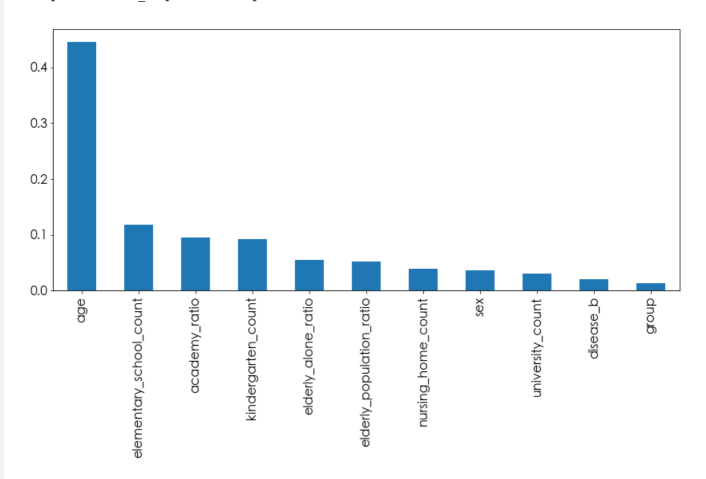
从下面的计算和作图中,我们可以看到年龄是最重要的特征,相比年轻人,高龄病人更容易死亡,我们可以看到文化水平和医疗水平等相关的变量的重要性远 小于年龄,而性别,是否为群体感染,是否患病对于病人是否死亡并没有特别大的作用。我们接下来对随机森林进行参数的调优。

### In [156]:

```
feature_imp = pd.Series(rf.feature_importances_)
feature_imp.index = x.columns
sorteds_features = feature_imp.sort_values(ascending=False)
sorteds_features
# The following is DataFrame format of the feature importance vector
feature_impDF = pd.DataFrame()
feature_impDF["importance"] = rf.feature_importances_
feature_impDF["feature"] = x.columns
feature_impDF.sort_values("importance",ascending=False,inplace=True)
feature_impDF
sorteds_features.plot(kind="bar", fontsize=16)
```

#### Out[156]:

<matplotlib.axes. subplots.AxesSubplot at 0x7f8ef5479710>



# 随机森林进行调优

## In [157]:

```
from sklearn.model_selection import GridSearchCV
import numpy as np
rf2 = RandomForestClassifier()
parameters = {"n_estimators":np.arange(10,100,10), "criterion":["gini", "entropy"]}
grid_rf2 = GridSearchCV(estimator=rf2, param_grid=parameters, cv=5,scoring="accuracy")
grid_rf2.fit(x_train,y_train)
```

#### Out[157]:

#### In [158]:

```
print(grid_rf2.best_params_)
```

{'criterion': 'gini', 'n\_estimators': 60}

根据网格计算,我们得到,最优的迭代为20次,此时的标准是gini指数。下面是运用这两个优化的参数进行新模型的建立。

# In [159]:

```
rf3 = RandomForestClassifier(class_weight='balanced',random_state=1,criterion="gini", n_estimators = 20)
###返回由训练集训练成的模型对验证集预测的结果
test_pred = rf3.fit(x_train,y_train).predict(x_test)
###打印混淆矩阵
print('\n'+'混淆矩阵: ')
```

```
print(con(y_test,test_pred))
###打印测试准确率
print('\n'+'Accuracy:')
print(rf3.score(x_test,y_test))

混淆矩阵:
[[ 13     4]
    [ 4 338]]

Accuracy:
0.9777158774373259
```

#### In [160]:

```
print(metrics.classification_report(y_test,test_pred))
```

|              | precision | recall | f1-score | support |
|--------------|-----------|--------|----------|---------|
| deceased     | 0.76      | 0.76   | 0.76     | 17      |
| released     | 0.99      | 0.99   | 0.99     | 342     |
| accuracy     |           |        | 0.98     | 359     |
| macro avg    | 0.88      | 0.88   | 0.88     | 359     |
| weighted avg | 0.98      | 0.98   | 0.98     | 359     |

优化后的新随机森林模型对于死亡病人的预测召回率有所下降,而对于康复病人的预测召回率有所提升,总体来讲,rg3模型对于病人的状态分类效果有所提升。 以下我们考虑 KNN 模型。

#### KNN

#### In [163]:

```
from sklearn.neighbors import KNeighborsClassifier
knn_model = KNeighborsClassifier(n_neighbors = 11)
knn_model.fit(x_train,y_train)
test_pred = knn_model.predict(x_test)
from sklearn import metrics
print(metrics.classification_report(y_test,test_pred))
print(metrics.confusion_matrix(y_test,test_pred))
print(metrics.accuracy_score(y_test,test_pred))
```

|               | precision | recall | f1-score | support |
|---------------|-----------|--------|----------|---------|
| deceased      | 0.62      | 0.47   | 0.53     | 17      |
| released      | 0.97      | 0.99   | 0.98     | 342     |
| accuracy      |           |        | 0.96     | 359     |
| macro avg     | 0.79      | 0.73   | 0.76     | 359     |
| weighted avg  | 0.96      | 0.96   | 0.96     | 359     |
| [[ 8 9]       |           |        |          |         |
| [ 5 337]]     |           |        |          |         |
| 0.96100278551 | 53204     |        |          |         |

我们可以看到,过半死亡状态的病人被预测为出院状态,因此对于死亡状态的预测不够准确,需要优化KNN模型。我们设置了k=1至21之间的奇数,对这些k的值都进行了模型拟合。

# In [164]:

```
k_list=[1,3,5,7,9,11,13,15,17,19,21]
for k in k_list:
    knn_model = KNeighborsClassifier(n_neighbors = k)
    knn_model.fit(x_train,y_train)
    test_pred = knn_model.predict(x_test)
    accuracy = metrics.accuracy_score(y_test,test_pred)
    confusion_mat = metrics.confusion_matrix(y_test,test_pred)

print("k = ",k)
```

```
print("\t正确率: ", '%.2f'%(accuracy*100) + "%")
   print("\t假阴性: ",confusion mat[0,1]/(confusion mat[0,1]+confusion mat[0,0]))
   print("","\t假阳性: ",confusion mat[1,0]/(confusion mat[1,0]+confusion mat[1,1]))
k = 1
 正确率:
       97.21%
 假阴性: 0.35294117647058826
 假阳性: 0.011695906432748537
k = 3
正确率: 97.77%
 假阴性: 0.35294117647058826
 假阳性: 0.005847953216374269
k = 5
 正确率: 96.66%
 假阴性: 0.5882352941176471
 假阳性: 0.005847953216374269
 正确率: 96.66%
 假阴性: 0.5882352941176471
 假阳性: 0.005847953216374269
k = 9
 正确率: 96.94%
 假阴性: 0.4117647058823529
 假阳性: 0.011695906432748537
k = 11
 正确率: 96.10%
 假阴性: 0.5294117647058824
 假阳性: 0.014619883040935672
k = 13
正确率: 95.82%
 假阴性: 0.7647058823529411
 假阳性: 0.005847953216374269
k = 15
 正确率: 95.82%
 假阴性: 0.7647058823529411
 假阳性: 0.005847953216374269
k = 17
 正确率: 96.10%
 假阴性: 0.7647058823529411
 假阳性: 0.0029239766081871343
k = 19
 正确率: 96.38%
 假阴性: 0.7058823529411765
 假阳性: 0.0029239766081871343
k = 21
 正确率: 96.38%
 假阴性: 0.7058823529411765
 假阳性: 0.0029239766081871343
```

可以看出预测出的假阴性所占的比例随k值增大逐渐上升,准确率逐渐下降,当k等于1的时候,假阴性为0.35,对于我们之前优化过的随机森林模型(假阴性为0.24),knn模型在这里并不是最优的模型。 综上我们选择调优过后的随机森林模型rg3作为一个较好的分类出院和病死的分类模型。

# Task3: 城市危险等级

目前仅对城市相关的数据进行了聚合,未想出如何去解释城市的聚类中心,以及如何设定适度函数。

## In [11]:

```
# task 3 risk levels of the cities
cr = covid_model.region.copy()
cr["count"] = 0
cr["released"] = 0
cr["isolated"] = 0
cr["deceased"] = 0

cnt = covid_model.patient_info.groupby(["province", "city"]).agg({'patient_id': ['count']})
cnt.columns = ['count']
cnt["released"] = 0
cnt["isolated"] = 0
cnt["isolated"] = 0
cnt["deceased"] = 0
cnt["deceased"] = 0
cnt = cnt.reset_index()
nc = pd.DataFrame(covid model.patient info.groupby(["province", "city"])["state"].value counts())
```

```
nc.columns = ['count']
nc = nc.reset index()
for row in nc.itertuples():
    cnt.loc[(cnt["province"]==getattr(row, "province"))&(cnt["city"]==getattr(row, "city")), getatt
r(row, "state")] = getattr(row, "count")
ccc = 0
for row in cnt.itertuples():
    if cr.loc[(cr["province"]==getattr(row, "province"))&(cr["city"]==getattr(row, "city"))].shape[
01 == 0:
        continue
    cr.loc[(cr["province"]==getattr(row, "province"))&(cr["city"]==getattr(row, "city")), "count"]
= getattr(row, "count")
    covid_model.region.loc[(covid_model.region["province"] == getattr(row, "province"))&(cr["city"] ==
getattr(row, "city")), "released"] = getattr(row, "released")
   cr.loc[(cr["province"]==getattr(row, "province"))&(cr["city"]==getattr(row, "city")), "isolated
"] = getattr(row, "isolated")
    cr.loc[(cr["province"]==getattr(row, "province"))&(cr["city"]==getattr(row, "city")), "deceased
"] = getattr(row, "deceased")
dcr = cr.drop(columns=["province", "city"])
k = 5
km = KMeans(n_clusters=k)
km.fit(dcr)
cent = km.cluster_centers_
km.labels
Out[11]:
array([2, 3, 4, 4, 4, 4, 4, 4, 0, 4, 0, 4, 4, 4, 4, 4, 4, 4, 4, 4, 4, 0,
       4, 4, 4, 4, 3, 0, 0, 0, 0, 0, 4, 4, 0, 0, 0, 0, 0, 0, 0, 0, 4, 3,
       0, 4, 0, 4, 4, 0, 4, 4, 3, 4, 0, 0, 4, 4, 3, 0, 0, 4, 4, 0, 4, 4,
       0, 0, 0, 3, 0, 0, 4, 4, 0, 4, 4, 0, 0, 0, 0, 0, 2, 0, 4, 0, 0, 0,
       0, 0, 4, 4, 0, 4, 4, 4, 4, 4, 0, 4, 0, 0, 0, 0, 0, 4, 0, 4, 0, 4,
       4, 0, 0, 4, 3, 0, 0, 0, 0, 0, 0, 0, 4, 0, 0, 0, 0, 0, 0, 0, 0,
       0, 3, 0, 0, 0, 0, 0, 0, 0, 0, 4, 0, 3, 0, 0, 0, 0, 0, 0, 0,
```

# Task4:基于地理的分析

4, 1], dtype=int32)

在附件中 task4.gif 这个动图展示了随着时间变化,病患往返路径的累计描述以及,病患活动地点的聚类中心的变化。聚类过程使用了 K-means 算法,并设定 k=10。可见病患活跃城市从一开始的很跳跃的状态逐步趋于稳定,中心皆为韩国人口众多的大城市。且从图中可以得出,一是韩国首都首尔连接的路径呈放射状,然后还有三至四条很明显的患病移动"干路"。切断这些城市之间的交通,可以比较好的延缓疫情发展态势。

#### In [167]:

```
# task 4 line route in
# define the kr map

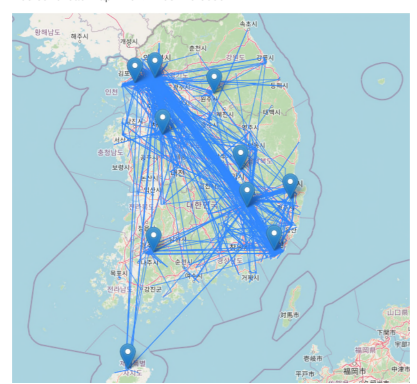
kkk = 10

loc = [35.8268093, 127.8512681]
kr_map = folium.Map(location=loc, zoom_start=7)

for k0, group in covid_model.patient_route.groupby("patient_id"):
    rt = list(zip(group["latitude"], group["longitude"]))
    folium.PolyLine(rt, weight=1).add_to(kr_map)

pos = covid_model.patient_route[["latitude", "longitude"]]
km = KMeans(n_clusters=kkk)
km.fit(pos)
cent = km.cluster_centers_
for point in range(0, kkk):
    folium.Marker(cent[point]).add_to(kr_map)

kr_map
```



# In [168]:

```
import datetime
import time
import imageio
import os,sys
from selenium import webdriver
from pprint import pprint
import glob
e=sys.exit
kkk = 10
swc = False
if swc:
    print(covid_model.patient_route["date"].max())
    begin = datetime.date(2020,2,9)
    end = datetime.date(2020,3,30)
    delta = datetime.timedelta(days=1)
    day = begin
    dest_folder = "tmp/route_gif/"
    while day <= end:</pre>
        loc = [35.8268093, 127.8512681]
        kr map = folium.Map(location=loc, zoom_start=8)
        ds = day.strftime("%Y-%m-%d")
        c day route = covid model.patient route.loc[covid model.patient route["date"] <= ds]</pre>
        if c_day_route.shape[0]:
            for k0, group in c_day_route.groupby("patient_id"):
                rt = list(zip(group["latitude"], group["longitude"]))
                folium.PolyLine(rt, weight=1).add_to(kr_map)
        pos = c day route[["latitude", "longitude"]]
        km = KMeans(n_clusters=kkk)
        km.fit(pos)
        cent = km.cluster_centers_
        for point in range(0, kkk):
            folium.Marker(cent[point]).add_to(kr_map)
        filename = ds+".html"
        kr map.save(dest folder+filename)
        folium map to png(filename, "/" + dest folder)
        # display(kr_map)
        # time.sleep(4)
        day += delta
if swc:
    images = []
    begin = datetime.date(2020,1,24)
    end = datetime.date(2020,3,30)
    delta = datetime.timedelta(days=1)
```

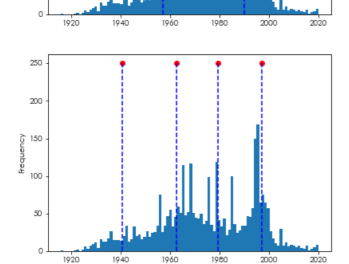
```
day = begin
dest_folder = "tmp/route_gif/"
duration = 10
while day <= end:
    filename = dest_folder + day.strftime("%Y-%m-%d") + ".png"
    images.append(imageio.imread(filename))
    day += delta
output_file = '1Gif.gif'
imageio.mimsave(dest_folder + output_file, images, duration=1/15)</pre>
```

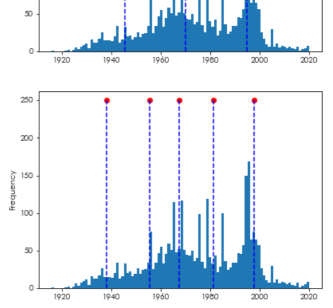
# Task6: 年龄分组

50

使用了 K-means 聚类算法, 分布展示了 k=2,3,4,5 时的年龄聚集中心。

```
In [171]:
# task 6
from sklearn.cluster import KMeans
kkk = 3
plt.figure()
plt.rcParams["figure.figsize"] = (14.0, 10.0)
for kkk in range(2, 6):
    birth = covid_model.patient_info["birth_year"]
    birth_np = np.array(birth)
    birth_np = birth_np.reshape(-1, 1)
    km = KMeans(n_clusters=kkk)
    km.fit(birth_np)
    cent = km.cluster_centers_
    # print(cent)
    plt.subplot(219 + kkk)
    ppt = covid model.patient info["birth year"].plot.hist(bins=100)
    plt.scatter(km.cluster_centers_, [250 for _ in range(kkk)], c='r')
    for i in range(kkk):
        plt.plot([cent[i], cent[i]], [0, 250], c='b', linestyle='--')
  250
                                                        250
  200
                                                        200
Fednency
100
                                                        150
  100
                                                        100
```





# SIR模型预测患病治愈趋势

对于Time.csv中每日累计患病,治愈,死亡的数据进行处理,得到每日实时患病,出院,死亡人数,结合每日全国的气温中位数temperature(简称tem)以及市民对于新冠肺炎的搜索趋势trend,进行SIR模型的建立:

```
dS/dt= -alS/N -b tem IS/N -c tem IS/N
dl/dt= alS/N +b tem IS/N +c tem IS/N - r1 I -r2 I
dR/dt= r1* I
dD/dt= r2* I
S:实时易感人群
I:实时感染人数
R:实时治愈人数
D:实时死亡人数
N:全国总人口,51269185
(a+b tem+c tem): 病毒传播的速率
在数据拟合中, 我们对其进行转化:
S(n+1) - S(n) = -aI(n)S(n)/N -b tem(n) I(n)S(n)/N -c tem(n) I(n)S(n)/N
I(n+1) - I(n) = aI(n)S(n)/N + b tem(n) I(n)S(n)/N + c tem(n) I(n)S(n)/N - r1 I(n) -r2 I(n)
R(n+1)-R(n)=r1*I(n)
D(n+1)-D(n)=r2*I(n)
I(n)表示第n天的实时感染人数。通过最小二乘法我们对参数 a, b, c, r1, r2进行估计,并对实时感染人数进行了拟合。
```

#### In [2]:

```
time=covid model.time
weatherdata=pd.read csv("../coronavirusdataset/Weather.csv")
search=pd.read_csv("../coronavirusdataset/SearchTrend.csv")
trend=search[1480:1558]["coronavirus"]/100
N=51269185
I=time["confirmed"]-(time["released"]+time["deceased"])
I=I.values
I=I[0:len(I)-1]
weather=weatherdata[23679:24927]["avg temp"].values
templist=[]
for i in range(78):
   sublist=[]
    for j in range(i*16,(i+1)*16):
        sublist.append(weather[j])
       mid=np.median(sublist)
   templist.append(mid)###77
tem=templist
del tem[-1]
R=time["released"]
R=R.values
R=R[0:len(R)-1]
D=time["deceased"]
D=D.values
D=D[0:len(D)-1]
RR=R+D
triS=N-I-RR
def delta(a):
   delta = [] #生成一个空列表,用来放新列表
    for i in range(len(a)-1):
        b = a[i+1] - a[i]
        delta.append(b)
    return delta
deltaSS=delta(triS)
deltaR=delta(R)
deltaD=delta(D)
deltaI=delta(I)
II=I[0:len(I)-1]
SS=triS[0:len(triS)-1]
```

```
tren=trena[U:1en(trena)-1]
col1=-II*SS/N
col11=col1.tolist()
col11.extend([i * (-1) for i in col11])
for i in [0]*154:
    coll1.append(i)
col1=col11
col2=-np.array(tem)*II*SS/N
col22=col2.tolist()
col22.extend([i * (-1) for i in col22])
for i in [0]*154:
    col22.append(i)
col2=col22
col3=-(tren**3)*II*SS/N
col33=col3.tolist()
col33.extend([i * (-1) for i in col33])
for i in [0]*154:
    col33.append(i)
col3=col33
col4=[0]*77
col4.extend([i for i in (-II).tolist()])
col4.extend([i for i in II.tolist()])
col4.extend([0]*77)
co15=[0]*77
col5.extend([i for i in (-II).tolist()])
col5.extend([0]*77)
col5.extend([i for i in II.tolist()])
X=pd.DataFrame({"col1":col1,"col2":col2,"col3":col3,"col4":col4,"col5":col5})
Y=deltaSS
Y.extend(deltaI)
Y.extend(deltaR)
Y.extend(deltaD)
Y=np.array(Y)
n=len(col1)
linreg = LinearRegression()
linreg.fit(X, Y)
print(linreg.coef )
print(linreg.score(X,Y))
[ 5.16939625e-02 -4.10907786e-03 8.09144275e-01 3.02697559e-02
```

```
7.76680308e-04]
0.7964748114306974
```

我们可以得到,参数a,b,c,r1,r2 分别是5.16939625e-02, -4.10907786e-03, 8.09144275e-01, 3.02697559e-02, 7.76680308e-04

下面我们对实时感染人数拟合,以及对治愈加上死亡人数的总和进行拟合,并作出趋势图。与实际数据相比,可以得出此模型拟合的效果较好。

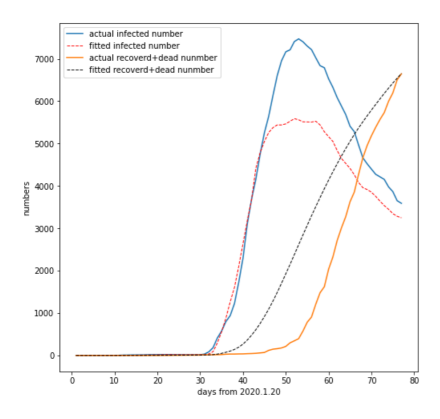
### In [3]:

```
a=0.0516939625
b=-4.10907786*0.001
c = 8.09144275*0.1
r1=3.02697559*0.01
r2=7.76680308*0.0001
ddI=a*II*SS/N+b*(tem*II*SS/N)+c*((tren**3)*II*SS/N)-r1*II-r2*II
ddR=r1*II
ddD=r2*II
ESI=1+ddI.cumsum()
ESR=ddR.cumsum()
ESD=ddD.cumsum()
xx=np.linspace(1,len(ESI),77)
plt.figure(figsize=(8, 8))
11=plt.plot(xx,II,label="actual infected number")
12=plt.plot(xx,ESI, color='red', linewidth=1.0, linestyle='--', label="fitted infected number")
13=plt.plot(xx,RR[0:len(RR)-1],label="actual recoverd+dead nunmber")
```

```
14=plt.plot(xx,(ESR+ESD), color='black', linewidth=1.0, linestyle='--',label="fitted recoverd+dead
nunmber")
plt.xlabel('days from 2020.1.20')
plt.ylabel('numbers')
plt.legend( loc='best')
```

# Out[3]:

<matplotlib.legend.Legend at 0x7ff9dfc6bb50>



# Conclusion

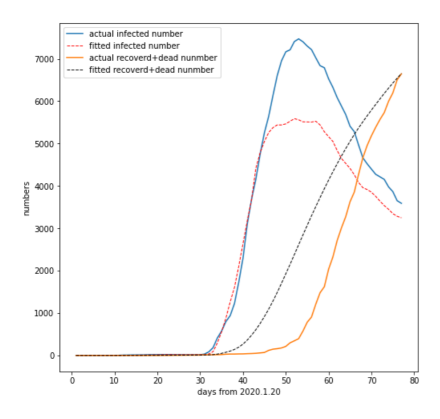
根据分析,我们发现,病人的年龄对于病人是否会死亡具有极大的影响作用。该项肺炎对于具有较低免疫力的高龄的人口(60代以上)具有较大的威胁,因此老人需要格外注意防护和隔离。由于韩国初期没有进行即时的防疫措施,最开始有病人在首尔周边以及大邱周边进行移动造成了病毒的传播,导致病毒在各个城市逐渐进行蔓延,最终达到急速增长的情况,因此在病毒发生初期即时地切断城市之间的联通渠道能够较好地防止病毒在全国范围内的广泛传播。此外我们还发现不同年龄段的人针对病毒的免疫力可以分成不同的类,较为高龄的人群更容易感染病毒。在四月下旬,韩国也颁布了法规,进行了相应的强制隔离措施,人们的防疫意识也有所提高。我们有理由相信韩国的未来疫情将会得到良好的改善。

In [ ]:

```
14=plt.plot(xx,(ESR+ESD), color='black', linewidth=1.0, linestyle='--',label="fitted recoverd+dead
nunmber")
plt.xlabel('days from 2020.1.20')
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plt.legend( loc='best')
```

# Out[3]:

<matplotlib.legend.Legend at 0x7ff9dfc6bb50>



# Conclusion

根据分析,我们发现,病人的年龄对于病人是否会死亡具有极大的影响作用。该项肺炎对于具有较低免疫力的高龄的人口(60代以上)具有较大的威胁,因此老人需要格外注意防护和隔离。由于韩国初期没有进行即时的防疫措施,最开始有病人在首尔周边以及大邱周边进行移动造成了病毒的传播,导致病毒在各个城市逐渐进行蔓延,最终达到急速增长的情况,因此在病毒发生初期即时地切断城市之间的联通渠道能够较好地防止病毒在全国范围内的广泛传播。此外我们还发现不同年龄段的人针对病毒的免疫力可以分成不同的类,较为高龄的人群更容易感染病毒。在四月下旬,韩国也颁布了法规,进行了相应的强制隔离措施,人们的防疫意识也有所提高。我们有理由相信韩国的未来疫情将会得到良好的改善。

In [ ]: