Air pollution and hospitalization of patients with idiopathic pulmonary fibrosis in Beijing: a time-series study

Lirong Liang1†, Yutong Cai2†, Baolei Lyu3, Di Zhang4, Shuilian Chu1, Hang Jing1, Kazem Rahimi4 and Zhaohui Tong1,5*

Abstract

Background: A small number of studies suggested that air pollution was associated with idiopathic pulmonary fibrosis (IPF) exacerbation, incidence and mortality. However, no studies to date were conducted in regions where air pollution is substantial. We aimed to investigate whether there are associations between acute increases in air pollution and hospitalization of patients with a confirmed primary diagnosis of IPF in Beijing.

Methods: Daily count of IPF hospitalizations (International Classification of Disease-10th Revision, J84.1) was obtained from an administrative database for 2013–2017 while daily city-wide average concentrations of PM10, PM2.5, NO2, Ozone, SO2 were obtained from 35 municipal monitoring stations for the same period. The association between daily IPF hospitalization and average concentration of each pollutant was analyzed with a generalized additive model estimating Poisson distribution.

Results: Daily 24-h mean PM2.5 concentration during 2013–2017 was 76.7 μg/m3. The relative risk (RR) of IPF hospitalization per interquartile range (IQR) higher (72 μg/m3) in PM2.5 was 1.049 (95% CI 1.024–1.074) and 1.031 (95% CI 1.007–1.056) for lag0 and moving averages 0–1 days respectively. No significant associations were observed for other lags. Statistically significant positive associations were also observed at lag0 with SO2, Ozone and NO2 (in men only). Positive associations were seen at moving averages 0–30 days for PM10 (RR per 86 μg/m3: 1.021, 95% CI 0.994–1.049), NO2 (RR per 30 μg/m3: 1.029, 95% CI 0.999–1.060), and SO2 (RR per 15 μg/m3: 1.060 (95% CI 1.025–1.097), but not with PM2.5 or Ozone.

Conclusions: Despite improvement in air quality since the implementation of clean air policy in 2013, acute exposure to higher levels of air pollution is significantly associated with IPF hospitalization in Beijing. Air quality policy should be continuously enforced to protect vulnerable IPF populations as well as the general public.

Keywords: Air pollutants, Interstitial lung diseases, Disease progression, Developing countries, Particulate matter

Background

Idiopathic pulmonary fibrosis (IPF) is the most common type of interstitial lung diseases (ILD), characterized by inflammation and fibrosis of the pulmonary parenchyma [1]. The exact causes of IPF are still largely unknown, with some studies suggesting that inhaled exposures such as cigarette smoke or wood dust may be among the triggers [2]. Patients with IPF experience
progressive deterioration of lung function, impaired quality of life and an estimated median survival of three to five years after diagnosis [1, 3, 4]. An acute respiratory worsening could significantly alter the natural history of IPF, potentially resulting in hospitalization and mortality. However, an immediate cause cannot always be established for around half of the IPF acute worsening cases [5].

Unlike major respiratory conditions (e.g. chronic obstructive pulmonary disease (COPD), bronchitis, asthma), for which evidence of harmful effects from ambient air pollution is well documented [6–8], evidence for IPF morbidity and mortality remains relatively scarce. Increased exposure to Ozone (O₃) were found to be associated with acute exacerbation of IPF six weeks after exposure in 436 and 192 patients in South Korea [5] and France [9] respectively, while number of hospitalizations was significantly higher following the days when levels of air pollutants including nitrogen dioxide (NO₂), Particulate Matter with a diameter less than or equal to 10 μm (PM₁₀) or 2.5 μm (PM₂.₅) were high in Santiago, Chile [10]. A more recent study of 152 patients in Japan concluded that exposure to PM₂.₅ and nitric oxide in the previous 30 days were significantly associated with IPF acute exacerbation [11]. Two studies in the United States found that higher air pollution exposure was associated with a lower level (i.e. the first phase of CAAP), and ran time-series analyses to investigate the acute effects of ambient air pollution on IPF hospitalization risk in Beijing. We additionally investigated whether the associations differed by patients’ characteristics, including age, sex and respiratory co-morbidity (COPD and asthma), as well as by seasonality.

Methods

Air quality data
Since 2013, major air pollutants have been routinely measured at 35 monitoring stations across Beijing, following the new Chinese national standard (GB 3095-2012). These monitoring stations were set up at locations that best represent different sources, for example, sources from road vehicles, urban anthropogenic activities, rural background and wider regional transport or background sources (Additional file 1: Fig. S1). At each station, a daily 24-h mean concentration for each pollutant except O₃ is reported based on hourly data that are available for at least 20 h in every 24 h. For O₃, a daily maximum 8-h moving mean concentration is reported based on hourly data that are available for at least 6 h in every 8 h.

Daily city-wide mean concentrations of PM₁₀, PM₂.₅, NO₂, SO₂ (sulphur dioxide), and O₃ were obtained from Jan 18th, 2013 to Dec 31st, 2017 (1809 days) by averaging daily mean readings from all 35 stations. Of the 1809 days, five days with missing data on all air pollutants were excluded. In addition, for O₃, daily mean concentration was missing for another 34 days. As a result, the analyses of PM₁₀, PM₂.₅, NO₂ and SO₂ were based on the 1804-day dataset whilst the analysis of O₃ was based on a 1770-day dataset. Daily mean temperature (°C) and relative humidity (%) were provided by the Beijing Meteorological Service website.

PM₂.₅, as a general air quality indicator, was not routinely monitored in Beijing before 2013. In order to examine whether there is difference on the association between PM₂.₅ and IPF hospitalization before and after the CAAP policy, we further obtained daily mean concentration of PM₂.₅ data during Apr 1st, 2008 and Dec 31st, 2012 (1508 days) from a single monitor operated by the US Embassy in Beijing. It was previously reported that within a radius of 40 km, this monitor covers 79.2% of the total population, 97.8% tertiary hospital and 79.3% secondary hospitals in Beijing that are eligible to admit IPF cases [19]. The PM₂.₅ data from this single monitor have been widely used in previous studies [19, 20]. To further verify the observations from this single PM₂.₅ monitor, we correlated the daily mean concentrations of this monitor with a nearby 1.5 km-apart government monitoring station using data from 2013. Overall, observations from the two monitors showed a high correlation (r = 0.93).

IPF hospitalization data
Daily counts of IPF hospitalization for the periods 2008–2012 and 2013–2017 were obtained from a database compiled by the Beijing Public Health Information Center. In Beijing, every government or private hospital at secondary or tertiary level is eligible to provide specialised care
and required to submit the front page of each electronic hospital discharge record to the database [18]. Detailed information including age, sex, residential address (at urban district or township level), admitted hospital, date of admission, primary discharge diagnosis and the corresponding International Classification of Diseases, 10th Revision (ICD-10) code was recorded. All hospitalization records for patients aged ≥18 years, resided in Beijing on a permanent basis (i.e. >6 months), and with a primary discharge diagnosis of IPF (ICD-10: J84.1) were extracted for this analysis. In Beijing, IPF patients could either self-refer to the outpatient department or be referred from primary care, and hospitalization will be warranted if their symptoms were found to be deteriorated. The primary diagnosis of IPF at discharge was coded by clinicians after reviewing the patient’s medical record (i.e. diagnostic terminology [10]) during hospitalization. All IPF admissions were from a total of 141 hospitals (75 tertiary and 66 secondary). Data were de-identified and it is not possible to identify individual patients, therefore informed consent was not required from this secondary data analysis. This study was approved by the Research Ethics Board of Beijing Chaoyang Hospital (approval number 2018-ke-303).

Statistical analyses
We defined same-day exposure as lag0 (e.g. the admission date) and examined a priori single-day exposure up to five days (single-day lag0 to lag4) and exposure of moving averages 0–1, 0–2, 0–3, and 0–4 before admission date.

While our primary analysis aim was to determine the acute effects of air pollution on IPF hospitalization, we additionally investigated the moving averages 0–30 period to account for potential subacute effects of air pollution on development of worsening symptoms of IPF. The exact exposure time window during which IPF patients develop an episode of worsening symptoms remains debated, although by convention the onset should be within one month prior to diagnosis [21]. Exposure time windows of some recent studies on the acute effects of air pollution on IPF hospitalization were confirmed IPF acute exacerbation cases were generally between the preceding 0–42 days [5, 11].

The associations between daily IPF hospitalization and average concentration of each pollutant were analyzed with a generalized additive model (GAM) estimating Poisson distribution:

\[
\text{Log}[E(Y_t)] = \text{intercept} + \beta C - i + ps(\text{calendar time}_t, 9) + ps(\text{temp}_t, 3) + ps(\text{RH}_t, 3) + \text{public holiday}_t + \text{day of week}_t
\]

where \(E(Y_t)\) represents the number of IPF cases on day \(t\); \(C\) is the city-averaged concentration of the pollutant and \(i\) is the day lag; \(\beta\) represents log-relative risk (RR) of IPF hospitalisation associated with a unit increase in the pollutant mean concentration; \(ps\) indicates penalised spline function to filter out long-term trends and seasonal patterns in daily IPF hospitalisations. \(\text{temp}\) and \(\text{RH}\) are the daily mean temperature (°C) and relative humidity (%) respectively at lag0; public holiday and day of week were included as categorical variables. Degrees of freedom (df) for calendar time, temperature and relative humidity were selected based on the parameters used in previous studies [22–24].

Each association was investigated in two-pollutant models if Spearman correlation ratios between these pollutants were less than 0.7. Subgroup analyses were conducted by age (18–64 years vs. ≥65 years), sex, season (warm vs. cool), asthma (yes vs. no) and COPD (yes vs. no). We conducted the sub-group analyses at lag0 only because for most studied pollutants, the association was found to be much stronger at lag0 in the main model. The Z-test was used to compare the two estimates derived from each sub-group.

The smoothing function of the generalized additive model was used to graphically analyse the exposure–response relationships between the log–RR of IPF hospitalization and air pollutant concentrations at lag0.

We did several sensitivity analyses by altering the generalized additive model: (1) to exclude calendar time, as long-term trends and seasonal patterns might also be partly related to pollutant concentrations; (2) to replace calendar time with an interaction term of exposure-by-season; (3) to increase the degrees of freedom of temperature and humidity to six; and (4) to model moving averages 0–4 of temperature and humidity instead of the current day (lag0). The latter two sensitivity analyses were to adjust potential non-linear and lagged confounding effects of weather conditions.

All statistical analyses were conducted in R (version 3.0.2) using packages MGCV, DPLYR, and TTR. Relative risk (RR) of hospitalisation for IPF per interquartile range (IQR) increase for each air pollutant were calculated and presented.

Results
Across Beijing, daily 24-h mean PM\(_{2.5}\) concentration during 2008–2012 and 2013–2017 was 88.9 and 76.7 μg/m\(^3\) respectively (Table 1), as compared to the recently updated WHO guideline of 15 μg/m\(^3\). For NO\(_2\) and SO\(_2\), daily 24-h mean concentration was 50.5 μg/m\(^3\) and 15.1 μg/m\(^3\) respectively; whilst daily 8-h mean concentration for O\(_3\) was 95.8 μg/m\(^3\). Spearman correlations between each pair of pollutants were low to moderate (i.e. r<0.7), except the high correlation between PM\(_{10}\) and PM\(_{2.5}\) (i.e. r>0.85). PM\(_{10}\) and PM\(_{2.5}\) have a permanent basis (i.e. >6 months), and with a primary discharge diagnosis of IPF (ICD-10: J84.1) were extracted for this analysis. In Beijing, IPF patients could either self-refer to the outpatient department or be referred from primary care, and hospitalization will be warranted if their symptoms were found to be deteriorated. The primary diagnosis of IPF at discharge was coded by clinicians after reviewing the patient’s medical record (i.e. diagnostic terminology [10]) during hospitalization. All IPF admissions were from a total of 141 hospitals (75 tertiary and 66 secondary). Data were de-identified and it is not possible to identify individual patients, therefore informed consent was not required from this secondary data analysis. This study was approved by the Research Ethics Board of Beijing Chaoyang Hospital (approval number 2018-ke-303).

Statistical analyses
We defined same-day exposure as lag0 (e.g. the admission date) and examined a priori single-day exposure up to five days (single-day lag0 to lag4) and exposure of moving averages 0–1, 0–2, 0–3, and 0–4 before admission date.

While our primary analysis aim was to determine the acute effects of air pollution on IPF hospitalization, we additionally investigated the moving averages 0–30 period to account for potential subacute effects of air pollution on development of worsening symptoms of IPF. The exact exposure time window during which IPF patients develop an episode of worsening symptoms remains debated, although by convention the onset should be within one month prior to diagnosis [21]. Exposure time windows of some recent studies on the acute effects of air pollution on IPF hospitalization were confirmed IPF acute exacerbation cases were generally between the preceding 0–42 days [5, 11].

The associations between daily IPF hospitalization and average concentration of each pollutant were analyzed with a generalized additive model (GAM) estimating Poisson distribution:

\[
\text{Log}[E(Y_t)] = \text{intercept} + \beta C - i + ps(\text{calendar time}_t, 9) + ps(\text{temp}_t, 3) + ps(\text{RH}_t, 3) + \text{public holiday}_t + \text{day of week}_t
\]

where \(E(Y_t)\) represents the number of IPF cases on day \(t\); \(C\) is the city-averaged concentration of the pollutant and \(i\) is the day lag; \(\beta\) represents log-relative risk (RR) of IPF hospitalisation associated with a unit increase in the pollutant mean concentration; \(ps\) indicates penalised spline function to filter out long-term trends and seasonal patterns in daily IPF hospitalisations. \(\text{temp}\) and \(\text{RH}\) are the daily mean temperature (°C) and relative humidity (%) respectively at lag0; public holiday and day of week were included as categorical variables. Degrees of freedom (df) for calendar time, temperature and relative humidity were selected based on the parameters used in previous studies [22–24].

Each association was investigated in two-pollutant models if Spearman correlation ratios between these pollutants were less than 0.7. Subgroup analyses were conducted by age (18–64 years vs. ≥65 years), sex, season (warm vs. cool), asthma (yes vs. no) and COPD (yes vs. no). We conducted the sub-group analyses at lag0 only because for most studied pollutants, the association was found to be much stronger at lag0 in the main model. The Z-test was used to compare the two estimates derived from each sub-group.

The smoothing function of the generalized additive model was used to graphically analyse the exposure–response relationships between the log–RR of IPF hospitalization and air pollutant concentrations at lag0.

We did several sensitivity analyses by altering the generalized additive model: (1) to exclude calendar time, as long-term trends and seasonal patterns might also be partly related to pollutant concentrations; (2) to replace calendar time with an interaction term of exposure-by-season; (3) to increase the degrees of freedom of temperature and humidity to six; and (4) to model moving averages 0–4 of temperature and humidity instead of the current day (lag0). The latter two sensitivity analyses were to adjust potential non-linear and lagged confounding effects of weather conditions.

All statistical analyses were conducted in R (version 3.0.2) using packages MGCV, DPLYR, and TTR. Relative risk (RR) of hospitalisation for IPF per interquartile range (IQR) increase for each air pollutant were calculated and presented.

Results
Across Beijing, daily 24-h mean PM\(_{2.5}\) concentration during 2008–2012 and 2013–2017 was 88.9 and 76.7 μg/m\(^3\) respectively (Table 1), as compared to the recently updated WHO guideline of 15 μg/m\(^3\). For NO\(_2\) and SO\(_2\), daily 24-h mean concentration was 50.5 μg/m\(^3\) and 15.1 μg/m\(^3\) respectively; whilst daily 8-h mean concentration for O\(_3\) was 95.8 μg/m\(^3\). Spearman correlations between each pair of pollutants were low to moderate (i.e. r<0.7), except the high correlation between PM\(_{10}\) and PM\(_{2.5}\) (i.e. r>0.85). PM\(_{10}\) and PM\(_{2.5}\) have
and PM2.5 ($r = 0.93$), PM10 and NO2 ($r = 0.73$), PM2.5 and NO2 ($r = 0.78$) (Additional file 1: Table S1). In 2013–17, 11,974 IPF admissions were recorded in Beijing, with a daily mean of six (Table 1). Number of admissions was higher among patients aged $\geq 65$ years, and patients without COPD, but no major differences were seen between sexes or seasons.

Acute exposure to higher PM2.5 exposure during 2013–17 was significantly associated with IPF hospitalization in a relatively short lag (Fig. 1, Additional file 1: Table S2). The RR per IQR (72 μg/m$^3$) higher was 1.049 (95% CI 1.024–1.074) at lag0, 1.031 (95% CI 1.007–1.056) at moving averages 0–1 and 1.021 (95% CI 0.998–1.044) at moving averages 0–2. Expressed per 10 μg/m$^3$ higher of PM2.5, the RRs are 1.007 (95% CI 1.003–1.010), 1.004 (95% CI 1.001–1.008) and 1.003 (95% CI 0.999–1.006) respectively. Results of PM10 were similar to those of PM2.5 except that for the moving averages 0–30, where there was a positive association with PM10 (RR: 1.021, 95% CI 0.994–1.049), compared to a null association with PM2.5 (RR: 0.994, 95% CI 0.962–1.027).

During 2008–2012, the RR of hospitalization for IPF at lag0 per interquartile range (IQR) (83 μg/m$^3$) higher in PM2.5 was 1.062 (95% CI 1.025–1.101) (Additional file 1: Fig. S2, Additional file 1: Table S3). The RR was reduced at lag1, lag2 and lag3 but remained positive. Positive associations with consistent RR estimates were observed for moving averages 0–1 up to 0–4. For example, the RR at moving averages 0–1 per IQR higher in PM2.5 was 1.054 (95% CI 1.028–1.090).

Associations varied by gaseous pollutants in different lags (Fig. 2, Additional file 1: Table S4). A marginally positive association was observed for each IQR (30 μg/m$^3$) higher of NO2 on both lag0 and moving averages 0–30 (RR: 1.029, 95% CI 0.999–1.060). A significant positive association was observed for SO2 at lag0 only (RR: 1.023 (95% CI 1.002–1.045) per IQR (15 μg/m$^3$) higher). Positive effect estimates of SO2 appeared to be in an increasing trend. For example, the RR at moving averages 0–1 and 0–30 was 1.019 (95% CI 0.996–1.043) and 1.060 (95% CI 1.025–1.097) respectively. A significant positive association was seen at lag0 for O3 (RR: 1.045, 95% CI 1.000–1.092 per IQR (85 μg/m$^3$) higher). Similar associations were observed at least until lag3. Significant positive associations were seen for moving averages 0–1 up to 0–4 (RR: 1.093, 95% CI 1.028–1.162); however, the association at moving averages 0–30 was positive but not statistically significant (RR: 1.047, 95% CI 0.941–1.165). The associations with O3 were statistically significant during cool season but not in warm season (Additional file 1: Table S5).

### Table 1: Daily air pollutant concentrations and IPF hospitalizations in Beijing, 2013–2017

| Air pollutant concentrations (μg/m$^3$) | Minimum | Maximum | Mean (SD) | Median (IQR) |
|---------------------------------------|---------|---------|-----------|--------------|
| PM10                                  | 0.0     | 220.0   | 109.7 (79.1) | 91.0 (54.0–140.0) |
| PM2.5 (2013–2017)                      | 0.0     | 467.0   | 76.7 (66.7) | 58.0 (29.0–101.0) |
| PM2.5 (2008–2012)                      | 0.0     | 552.5   | 88.9 (73.7) | 69.0 (36.0–118.9) |
| NO2                                   | 0.0     | 155.0   | 50.5 (24.2) | 44.0 (33.0–63.0) |
| SO2                                   | 0.0     | 139.0   | 15.1 (18.4) | 8.0 (4.0–19.0) |
| O3                                    | 0.0     | 292.0   | 95.8 (62.2) | 83.0 (50.0–135.0) |
| O3, cool season                        | 0.0     | 292.0   | 128.5 (63.5) | 126.0 (78.0–175.0) |

IPF hospitalization (number of cases per day), 2013–2017 (n = 11,974)

|                         | Minimum | Maximum | Mean (SD) | Median (IQR) |
|-------------------------|---------|---------|-----------|--------------|
| Total                   | 0       | 22      | 6.4 (3.9) | 6.0 (3.0–9.0) |
| Male                    | 0       | 14      | 3.6 (2.5) | 3.0 (2.0–5.0) |
| Female                  | 0       | 12      | 2.8 (2.1) | 2.0 (1.0–4.0) |
| Age < 65 years          | 0       | 12      | 1.9 (1.8) | 2.0 (1.0–3.0) |
| Age $\geq$ 65 years     | 0       | 17      | 4.5 (2.9) | 4.0 (2.0–6.0) |
| Warm season, May to October | 0   | 22      | 6.4 (3.8) | 6.0 (3.0–9.0) |
| Cool season, November to April | 0   | 21      | 6.5 (4.0) | 6.0 (3.0–9.0) |
| With COPD*              | 0       | 4       | 0.4 (0.7) | 0.0 (0.0–1.0) |
| Without COPD            | 0       | 21      | 6.0 (3.7) | 6.0 (3.0–9.0) |
| With asthma             | 0       | 2       | 0.1 (0.3) | 0.0 (0.0–0.0) |
| Without asthma          | 0       | 21      | 6.4 (3.9) | 6.0 (3.0–9.0) |

* COPD chronic obstructive pulmonary disease

COPD chronic obstructive pulmonary disease
The main results remained stable in all the sensitivity analyses (Additional file 1: Table S6) and two-pollutant models (Additional file 1: Table S7). During 2013–2017, positive exposure–response curves at lag0 were observed at least up to 300 μg/m^3 for PM_{2.5}, whereas various patterns were observed for the gaseous pollutants (Additional file 1: Fig. S3).

Stronger positive associations with all pollutants except O_3 were seen in men as compared to women, with the interaction term with NO_2 reaching statistical significance (Fig. 3). In men, per IQR higher of exposure to NO_2 at lag0 was significantly associated with IPF hospitalization (RR: 1.064, 95% CI 1.023–1.107). While in women, per IQR higher exposure to O_3 at lag0 was significantly associated with IPF hospitalization (RR: 1.087, 95% CI 1.017–1.161). There were no significant effect modifications by age, season, COPD or asthma with each pollutant, although there is a tendency that patients aged ≥ 65 years had a higher hospitalization risk with higher exposure to each pollutant except NO_2. For IPF patients with COPD, hospitalization risk (RR: 1.230, 95% CI 1.030–1.469) was significantly higher as per IQR higher O_3 exposure, as compared to patients without COPD (RR: 1.034, 95% CI 0.988–1.082).

**Discussion**

Our findings suggest that acute exposure to higher levels of air pollutants, both particulates and gases, was significantly associated with IPF hospitalization in Beijing during 2013–2017. The magnitude of the associated risk is comparable to what was previously observed in Beijing for COPD exacerbation requiring hospitalization [18].

To date, only four studies reported associations between short-term air pollution exposure and acute exacerbation or hospitalization of IPF [5, 9–11], with mixed findings observed across the studies. This may be due to the differences in exposure levels and toxicity, study populations in different climate zones and ethnic groups, analytical approaches as well as the criteria in diagnosing and/or admitting IPF patients.

Our finding of an association between acute PM_{2.5} exposure and IPF hospitalization is an important addition to current knowledge. Previously, the Korean study did not have PM_{2.5} data [5], whilst the French study did
not find an association between short-term PM$_{2.5}$ and acute exacerbation of IPF among 192 patients [9]. The time-series study in Santiago, Chile reported that PM$_{2.5}$ exposure in the previous four days was a significant risk factor of IPF hospitalization [10], which is in line with our study. Despite the concentrations of ambient PM$_{2.5}$ was much higher in Beijing (77 μg/m$^3$ 2013–2017) than in Santiago (29 μg/m$^3$ 2001–2012), the risk estimate in our study was relatively smaller. This may reflect differences in statistical approaches used (e.g. the Chilean study ran separate region-specific time-series analyses and then pooled all the estimates together), PM$_{2.5}$ toxicity and population characteristics.

Our study has suggested that more acute exposure (i.e. past three days) to PM$_{2.5}$ as compared to more recent exposure (i.e. past 30 days), was relatively more important in leading to IPF hospitalization. This is in contrast to a Japanese study of 152 patients in which a significant positive association between IPF acute exacerbation and prior 30-day average PM$_{2.5}$ exposure was found [11]. We did find, however, a positive association between prior 30-day average PM$_{10}$ exposure and IPF hospitalization risk. This finding highlighted the potential of coarse PM in triggering symptoms among IPF patients via direct deposition in the respiratory tract over a sub-acute period [24]. In Beijing, studies have shown that sources of PM$_{10}$ mainly come from dust whilst for PM$_{2.5}$ traffic is the largest source with coal combustion and biomass burning contributing in winter months [25, 26]. We suspect that the differences in sources as well as compositions may have influenced the time course of both coarse PM and fine PM on IPF hospitalization. More research is needed to confirm this, especially in cities where sources of air pollution are diverse. Numerous mechanistic studies have indicated that PM could cause oxidative stress via over-production of reactive oxygen species [2, 24], which eventually leads to cellular damage and shortened telomeres, a recognized risk factor of IPF [27]. Both coarse and fine PM could also induce local and systemic inflammation of similar magnitude [28]. Collectively, these biological pathways could accelerate the lung injury and fibrotic process among the IPF patients.

We found that, as compared to the associations with PM$_{2.5}$ observed in 2013–2017, the associations observed in 2008–2012 were relatively stronger and the lag time to hospitalization was longer (5 vs. 2 days). It should be noted that PM$_{2.5}$ data from 2008 to 2012 were only based on a single monitor. Although this is not a direct assessment of health benefits resulting from the Clean Air Action Plan, our findings seem to echo previous reports that improvement in air quality has indeed improved several morbidity and mortality outcomes in the Chinese population [18, 29, 30].

This study found that acute exposure to O$_3$ up to preceding five days was significantly associated with IPF hospitalization. Further stratifying the associations by season, only in cool season can the statistically significant positive associations be observed. This is somehow unexpected as previous studies in Beijing or the northern part of China found that short-term effects of O$_3$ on COPD exacerbation [18] and daily mortality [31] were more pronounced in warm season due to the high ambient concentration, the possible synergistic effects with high temperature as well as the fact that people spend more time outdoors. Previously, Johannson et al. only analysed O$_3$ concentrations in summer months and reported a significant positive association with acute exacerbation of IPF [5] whilst Sesé et al. also reported a significant positive association, but without stratifying the risk by season [9]. The studies in Chile [10] and Japan [11] did not report an association with O$_3$. As ours is the only study to date that reported seasonal variations of O$_3$ associations with IPF hospitalization, these patterns remain to be determined in future studies. An alternative explanation is that O$_3$ exposure in cool season may represent a proxy of other more important factors, such as viral infection, which is an known trigger in worsening IPF that may require hospitalization [4]. Being a strong oxidant, O$_3$ is
Fig. 3 Subgroup analyses on the associations between each pollutant (per IQR higher) and hospitalization for IPF in single-pollutant models at Lag0.
known to have the capability to exacerbate chronic lung conditions including asthma and COPD. In addition, O₃ can cause breathing difficulty by bronchoconstriction, thereby increasing the risk of hospitalization among individuals with existing respiratory diseases including IPF [32]. Animal and human studies have suggested that O₃ can reach and accumulate at the lower lungs because of its low water solubility, leading to inflammation and impaired small airways function, and potential hospitalization risk [33, 34].

Previous studies either found no associations between acute exacerbation of IPF and SO₂ exposure [5], or that the associations appeared to be confounded by PM [10]. Despite a remarkable reduction of SO₂ concentrations by 70% was recorded in Beijing from 2013 to 2017 [18], we still observed that IPF hospitalization risk was significantly higher following recent periods (i.e. past 30 days) of high ambient SO₂ concentration. Associations did not appear to be significantly affected by further adjustment of PM₂.₅. These results indicated that continued investment on renewable and clean energy, with a steady reduction of traditional fossil energy consumption, still remain pivotal to SO₂ abatement in order to mitigate the harmful impacts on respiratory diseases. As with O₃, patients with chronic respiratory diseases may be sensitive to the effects of SO₂, which has the capacity to induce oxidative stress, airway inflammation and breathing difficulty [35].

We observed a marginally positive association between concurrent-day NO₂ exposure and IPF hospitalization in the overall population. Interestingly, we found the same effect size for both moving averages 0–30 and lag0. Only in the men-only sample, this positive association became statistically significant. This finding is partly in line with the Johannson study of 436 (80% of them were men) South Korean patients [5], in which higher exposure to previous six-week average NO₂ was significantly associated with IPF exacerbation. It should be noted that, association with the most acute exposure period (i.e. the week before manifested IPF exacerbation) was weaker and not statistically significant in the Johannson study. Similarly, the Japanese study of 152 patients (70% of them were men) reported significant positive associations between prior 30-day average NO₂ and IPF acute exacerbation. In contrast, the French study did not find any association with NO₂ [9]. The time-series study in Chile observed robust positive associations between acute increased exposure to NO₂ in the previous four days and IPF hospitalization in all populations, with the association being stronger among women [10]. NO₂ has been commonly studied as an indicator of traffic emission and is usually highly correlated with PM₂.₅, therefore it remains difficult to study its independent effects. Nonetheless, some have argued that NO₂, via the airway inflammation pathway, could have direct and adverse impacts on respiratory health [36].

We observed that the risk of O₃ on IPF hospitalization was more pronounced in women, in line with studies which generally found women had a higher respiratory hospitalization risk from O₃ exposure, particularly during the cold season [33]. And for NO₂ effects, we only found significant associations in men. The sex-specific (biological factors, e.g. sex hormones, airway diameter, lung size) and gender-specific (socioeconomic and cultural factors, e.g. occupation, personal activity) differences of air pollution effects on respiratory health outcomes still largely remain inconsistent across studies [33], and future studies are warranted to gain better insights into these two different pathways.

To our best knowledge, this is by far the first and the largest time-series study that was conducted in a city with high levels of ambient air pollution, using a relatively large number of IPF hospitalizations over a 5-year period from a representative database. The study has limitations. First, it is an ecological study by design, and hence the modelled risk estimates should not be interpreted as predictive of individual hospitalisation probability. Given the time-series study design, we were unable to adjust for important individual-level factors of IPF such as smoking and socioeconomic status. However, the proportion of these risk factors are unlikely to change substantially in the same population which we were investigating over a short period of time. Notably, time-varying factors such as seasonal viral or bacterial infection will likely have a more important role to confound these relationships, but we did not have these data for further investigation. Second, the use of daily, outdoor, city-wide average air pollutant concentrations inevitably will have introduced bias to health estimates because we cannot take into account place of residence and time-activity patterns (e.g. workplace exposure, travel, indoor cooking) from the study population. Due to privacy reasons, we were only provided area-level (urban district or township) residential address for each patient for this secondary analysis, therefore we cannot model air pollution exposure at individual residence. However, this time-series analysis is an important first step to inform our next immediate work to study the impact of modelled air pollution at residence on health outcomes in a cohort of IPF patients should consent of accessing address data was granted. In addition, personal monitoring of air pollution exposure among IPF patients will provide even more novel insights into the exposure patterns and the associated health risks. Third, as with the Chilean study [10], our outcome was a primary diagnosis of IPF (ICD-10 J84.1) at discharge, which was coded by clinicians and usually the main reason for admission or length of stay. Most
IPF-related hospitalizations under J84.1 would be likely related to pneumonia or acute exacerbation, and it is true that neither of which typically manifest clinically within days of a triggering exposure. Therefore, cautious interpretation is needed on our observed significant positive associations in a relative short lag of higher air pollution exposure. Fourth, models including more than two pollutants were not possible due to the high correlation among pollutants. Also, we lacked the information on the occupational history of these patients. Lifetime occupational exposure might have altered the air pollution risk on respiratory health [7] and this is particularly relevant considering the rapid industrialization in Beijing in the last few decades. As pointed out already in the Chilean study, these time-series analyses were mainly to determine whether there was an association between air pollution and acute worsening of IPF that severe enough for hospitalisation. We, however, were unable to determine whether air pollution is the primary cause, or aggravates other intermediate disorders, that ultimately leads to rapid deterioration in IPF patients [10].

Conclusions

The implementation of Clean Air Action Plan in Beijing successfully reduced the annual mean concentration of PM$_{2.5}$ by one-third (from 87 μg/m$^3$ in 2013 to 58 μg/m$^3$ in 2017) in the first phase. Although the annual mean PM$_{2.5}$ in Beijing decreased to around 40 μg/m$^3$ by end of 2021, it is still eight times higher than the recently updated WHO target value of 5 μg/m$^3$, and therefore remains a significant health threat as demonstrated in this study. The concentrations for nitrogen dioxide and ozone, both of which are known to be harmful for respiratory health [31] and the climate, however, remain relatively stable, indicating stringent emission controls are needed, particularly on traffic which is the primary source in Beijing. Tackling these air pollutants should also be recognized as a key player in policies aiming for mitigating impacts from climate change, results of which will also bring health co-benefits.

In conclusion, this study provides evidence that acute exposure to higher levels of air pollution is associated with IPF hospitalization in Beijing. Local air quality policy should be stringently enforced, and carefully monitored for the progress to protect vulnerable populations, the general public and the planet.

Abbreviations

CAAP: Clean Air Action Plan; COPD: Chronic obstructive pulmonary disease; FVC: Forced vital capacity; GAM: Generalized additive model; ICD-10: International Classification of Diseases, 10$^{th}$ Revision; ILD: Interstitial lung diseases; IPF: Idiopathic pulmonary fibrosis; IQR: Inter-quartile range; NO$_2$: Nitrogen dioxide; O$_3$: Ozone; PM: Particulate matter; RR: Relative risk; SO$_2$: Sulphur dioxide; WHO: World Health Organisation.

Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1186/s12931-022-01998-8.

Additional file 1. Figure S1. Air quality monitoring stations across Beijing.

Table S1. Spearman correlation between air pollutants during 2013-2017.

Table S2. Associations between PM$_{2.5}$ and PM$_{10}$ (per IQR increase) and IPF hospitalisation in Beijing at different lags during 2013-2017.

Table S3. Associations between PM$_{2.5}$ (per IQR increase of 83 μg/m$^3$) and IPF hospitalisation in Beijing at different lags during 2008-2012.

Figure S2. Comparisons of associations between PM$_{2.5}$ and IPF hospitalization for periods 2008-2012 and 2013-2017.

Table S4. Associations between gaseous pollutants (per IQR increase) and IPF hospitalisation in Beijing at different lags during 2013-2017.

Table S5. Associations between ozone (per IQR increase) and IPF hospitalisation in Beijing at different lags during 2013-2017 by season.

Table S6. Results of the sensitivity analyses.

Table S7 Results of two-pollutant models. Figure S3. Exposure-response curves between air pollution exposure and IPF hospitalization at lag0.

Acknowledgements

We thank the data providers and their data collection teams.

Authors’ contributions

LL, YC and ZT conceived and designed the study. LL and YC led the statistical data analysis. BL acquired and provided inputs on the air quality data. DZ, SC and HJ acquired the health-related data. YC wrote the first draft of the manuscript, with all authors contributing to the critical revision of the manuscript for intellectual content. All authors reviewed and approved the final manuscript.

Funding

This research is funded by Beijing Municipal Science & Technology Commission (Grant number: Z201100005520028), and Beijing Municipal Administration of Hospitals Incubating Program (Grant number: PK2020014). YC and KR are supported by the PEAK-Urban Programme, funded by UK Research and Innovation’s Global Challenge Research Fund (Grant number: ES/P011055/1). YC acknowledges support from National Institute for Health Research (NIHR) Health Protection Research Unit in Environmental Exposures and Health, a partnership between the UK Health Security Agency, the Health and Safety Executive and the University of Leicester. The views expressed are those of the author(s) and not necessarily those of the NHS, the NIHR, the Department of Health and Social Care or the UK Health Security Agency.

Availability of data and materials

The hospitalisation datasets used and/or analysed for the current study was provided by Beijing Public Health Information Center but restrictions apply to the availability of these data, which were used under license for the current study, and so are not publicly available. Data are however available from the authors upon reasonable request and with permission of Beijing Public Health Information Center. Air quality data are publicly available.

Declarations

Ethics approval and consent to participate

This study was approved by the Research Ethics Board of Beijing Chaoyang Hospital (approval number 2018-ke-303).

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

Author details

1 Department of Clinical Epidemiology & Tobacco Dependence Treatment Research, Beijing Institute of Respiratory Medicine, Beijing Chaoyang Hospital, Capital Medical University, Beijing, China. 2 Centre for Environmental Health and Sustainability, Department of Health Sciences, University of Leicester,
