Severe volcanic SO$_2$ exposure and respiratory morbidity in the Icelandic population – a register study

Hanne Krage Carlsen (✉ hkc1@hi.is) 
University of Iceland  https://orcid.org/0000-0003-1656-9624

Unnur Valdimarsdóttir
School of public health sciences, University of Iceland

Haraldur Bríem
Chief Epidemiologist, Directorate of Health, Centre of Health Threats and Communicable Diseases

Francesca Dominici
Department of Epidemiology, Harvard T H Chan School of Public Health

Ragnhildur Gudrun Finnbjörnsdottir
The Environment Agency of Iceland

Thorsteinn Jóhannsson
The Environment Agency of Iceland

Thor Aspelund
School of Health Sciences, University of Iceland

Thorarinn Gislasson
Landspitali - The National University Hospital

Thorolfur Gudnason
Centre for Health Threats and Communicable Diseases, Directorate of Health

Research

Keywords: Volcanic eruption, atmospheric transport, respiratory disease, epidemiology, public health

Posted Date: December 1st, 2020

DOI: https://doi.org/10.21203/rs.3.rs-68431/v2

License: © This work is licensed under a Creative Commons Attribution 4.0 International License. Read Full License
Version of Record: A version of this preprint was published at Environmental Health on February 27th, 2021. See the published version at https://doi.org/10.1186/s12940-021-00698-y.
Abstract

Background

The Holuhraun volcanic eruption September 2014 to February 2015 emitted large amounts of sulfur dioxide (SO$_2$). The aim of this study was to determine the association between volcanic SO$_2$ gases on general population respiratory health some 250 km from the eruption site, in the Icelandic capital area.

Methods

Respiratory health outcomes were: asthma medication dispensing (AMD) from the Icelandic Medicines Register, medical doctor consultations in primary care (PCMD) and hospital emergency department visits (HED) in Reykjavík (population: 215 000) for respiratory disease from 1 January 2010 to 31 December 2014. The associations between daily counts of health events and daily mean SO$_2$ concentration and high SO$_2$ levels (24-hour mean SO$_2$>125µg/m$^3$) were analysed using generalized additive models.

Results

After the eruption began, AMD was higher than before (129.4 vs. 158.4 individuals per day, $p<0.05$). For PCMD and HED, there were no significant differences between the number of daily events before and after the eruption (142.2 vs 144.8 and 18.3 vs 17.5, respectively). In regression analysis adjusted for other pollutants, SO$_2$ was associated with estimated increases in AMD by 0.99% (95% CI 0.39 - 1.58%) per 10 µg/m$^3$ at lag 0-2, in PCMD for respiratory causes 1.26% (95% CI 0.72 - 1.80%) per 10 µg/m$^3$ SO$_2$ at lag 0-2, and in HED by 1.02% (95% CI 0.02- 2.03%) per 10 µg/m$^3$ SO$_2$ at lag 0-2. For days over the health limit, the estimated increases were 10.9% (95% CI 2.1-19.6%), 17.2% (95% CI 10.0-24.4%) for AMD and PCMD.

Dispensing of short-acting medication increased significantly by 1.09% (95% CI 0.49-1.70%), and PCMD for respiratory infections and asthma and COPD diagnoses and increased significantly by 1.12% (95% CI 0.54-1.71%) and 2.08% (1.13-3.04%).

Conclusion

High levels of volcanic SO$_2$ are associated with increases in dispensing of AMD, and health care utilization in primary and tertiary care. Individuals with prevalent respiratory disease may be particularly susceptible.

Introduction

SO$_2$ (Sulphur dioxide) exposure is associated with respiratory health morbidity and mortality(1, 2) and at higher concentrations (a 10 minute mean over 500 µg SO$_2$ per m$^3$) it is associated with irritation of the respiratory tract in susceptible individuals(2) and can trigger respiratory symptoms such as acute bronchial asthma, pulmonary oedema, and respiratory distress(3) – especially in individuals with hyper-reactivity syndrome.(4) Populations within 100 km of a volcanic eruption have traditionally been
considered at risk (5) although studies have found possible health effects of volcanic ash at greater distances (6).

During the Holuhraun volcanic eruption in the Barðarbunga central volcanic system SO$_2$ was dispersed widely over Iceland according to meteorological conditions, reaching the capital area some 250 km from the eruption site where the 24-hour air quality guideline limit for SO$_2$, $125 \mu g/m^3$ (2), was exceeded repeatedly during the fall of 2014. The Barðarbunga volcanic system is located in the central highlands of Iceland which is uninhabited and no humans live within 50 km of the eruption site so although there was no immediate danger, several steps were taken to inform and advise the public of the situation with the dispersed SO$_2$ gas. Press briefings and community meetings were held from mid-September and the civil protection agency issued warnings by text messages to all cell phones in the affected areas when high levels of SO$_2$ were expected. During the eruption, there were no formal disruptions to daily life such as school closings. (8)

The eruption in the fall and winter of 2014-2015 was the largest eruption in Iceland since the Laki eruption in 1783-1784. Some 12 million tons of sulphur dioxide, SO$_2$, was emitted from the eruption and the resulting lava field,(9) A clinical study of professionals working at the eruption site with very high exposures found no serious health effects associated with exposure, perhaps because they were wearing protective equipment, most importantly, masks.(10)

Exposure to SO$_2$ from active volcanoes is associated with increased rates of chronic cough and phlegm, as well dry and sore throat,(11-18) although few fatalities have been reported from very high SO$_2$ exposure near active volcanoes.(19) In Hawai‘i, PM$_{2.5}$ (particle matter with an aerodynamic diameter $<2.5$ µm) with a significant amount of volcanic emissions was associated with increased respiratory admissions.(20)

While concentration-response relationships between volcanic SO$_2$ and respiratory symptoms, many of the studies’ designs and methods leave them prone to bias. For example, symptoms are often self-reported, or participants were aware of their exposure status,(21) whereas other studies suffer from a lack of data. (22) Moreover, most of the existing literature pertains to long-term area-wide exposure whereas SO$_2$ exposure in Iceland during the Holuhraun eruption was intermittent with few hours or days of high SO$_2$ concentrations followed by periods of low SO$_2$ concentrations as wind directions changed.(7) With population-based registers on medicine dispensing and health care utilization as well as vigorous air pollution monitoring in the capital area, Iceland provides an ideal setting for studying population health effects of short-term exposure to SO$_2$ from a volcanic eruption.

The objective of this study was to study the acute effects of exposure to SO$_2$ concentrations from a volcanic source and SO$_2$ concentrations above the air quality guideline value of 24 hour mean of 125$\mu g/m^3$) on respiratory health in the general population and to investigate risk differences in subgroups of the population and susceptible groups.
Material And Methods

The Holuhraun volcanic eruption in North-East central Iceland began 31 August 2014 and ended 27 February 2015. The study period was 1 January 2010 – 31 December 2014, and the time before the eruption was used a reference period. The Holuhraun eruption persisted until end of February 2015, whereas our study period ends 31 December 2014 due to a change in the database recording of events. However, SO$_2$ never exceeded the 24-hour air quality guideline limit during January and February 2015, although daily mean concentrations were still higher than before the eruption.

The mean population of Iceland during the study period was 320 000 inhabitants. The capital area, Reykjavik and surrounding municipalities, had 205 282 residents at the beginning of the study period, and 215 965 residents at the end.(23) The analysis was restricted to the capital area (residential postcodes 101-171, 200-225, and 270) where adequate information about SO$_2$ exposure was available for the whole study period. The Icelandic health care system is state-centred, mainly publicly funded system with universal coverage.(24) We obtained data on respiratory health and individual data on residence (postcode), age, sex and an anonymous personal identification number from 1) the National Medicines Register; 2) Primary care centres (that function as first point of contact) and 3) Landspitali, the national university hospital, the country’s centre of clinical excellence.(2) All registers are held by the Icelandic Directorate of Health and extraction is subject to approval from the Icelandic Bioethical Committee. From the National Medicines Register we extracted data on dispensing (pharmacy sales to individuals) of prescription anti-asthma medication (AMD) classified by The World Health Organisation Anatomical Therapeutic Chemical code R03. AMD relieve symptoms of asthma and chronic obstructive pulmonary disease, COPD, and are occasionally prescribed to individuals with respiratory infections. Furthermore, AMD is a proxy for respiratory health in a population.(25-29) From the primary care centers (PCC) and hospital emergency department (HED) databases at the Directorate of Health we extracted data on individuals diagnosed with respiratory illnesses.

In the main analyses, we analysed the number of MD visits in primary care (PCMD) and all HED visits regardless of admission status. As the same bout of illness is likely to result in recurring contacts with the health care system, we included only the first record of an individual’s health care contacts within a 14-day period for the same diagnosis category to avoid exposure misclassification with respect to the timing of the outcome. For each outcome, we constructed daily time series starting 1 January 2010 to 31 December 2014 for the following age groups; children (under 18 years of age), adults (18-64 years), and elderly (age 65 years and above), see data selection in Flow diagrams 1-3 in the supplemental material. We obtained SO$_2$, PM$_{10}$ (particle matter with an aerodynamic diameter <10 µm), and NO$_2$ (nitrogen dioxide) data along with meteorological data from the Icelandic Environment Agency’s stationary air pollution monitor located in Reykjavik (Figure 1) for the study period and constructed a time series of 24-mean values from midnight to midnight.

Statistical methods
Descriptive statistics were calculated for all exposure and outcome variables for the period before and after the beginning of the eruption (for health data, we used age at date of first occurrence in the data) (see Table 1, see also supplementary material, Flow diagrams 1-3). Correlations of the exposure variables can be found in the supplement (Table S1). We use the t-tests to assess as whether there was a statistically significant difference in concentrations of relevant pollutants and the number of daily health outcome events before, and during the eruption (Table 1 and Table 2).

In the regression analysis of the daily number outcomes during the whole study period, SO\textsubscript{2} exposure was given as either a) a continuous variable, or b) an indicator value of the 24-hour SO\textsubscript{2} concentration exceeding the air quality guideline value (125 µg/m\textsuperscript{3}).

We fitted distributed lag non-linear models (DNLM) to the data. (30) to identify the delay in days (lag days) from exposure to the observed health outcomes (Supplemental Figure S2) and concentration response (Supplemental Figure S3 and Figure S4). We estimated the effects of SO\textsubscript{2} exposure on the outcome by fitting generalized additive models (GAM). (31)

\[ Y_t \sim \text{Quasipoisson} (\mu_t) \]

\[ \log \mu_t = \alpha + \beta_1 \text{SO}_2 + \beta_2 \text{PM}_{10} + \beta_3 \text{NO}_2 + \beta_4 \text{RelativeHumidity} + \beta_5 I_{dow} + \beta_6 \text{Strike} + \beta_7 Y_{t-1} \]

\[ + s_1(\text{Temperature}) + s_2(\text{Day in the time series}) + s_3(\text{Day of the year, } bs=\text{"cc"}) \]

Where \( Y_t \) denotes the daily number of health events, \( \beta_1 \) denotes the log relative rate of events associated with a 10 µg/m\textsuperscript{3} increase in SO\textsubscript{2} at lag 0-2. Furthermore, the results are adjusted for co-pollutants and weather (PM\textsubscript{10}, NO\textsubscript{2}, and relative humidity) at the same lag intervals as the main exposure. \( I_{dow} \) is an indicator for day of week and odd holidays, and Strike is an indicator of strike days (used only in analysis of the hospital and primary care data, to indicate days where medical doctors went on strike as part of a labor conflict, which coincided with some high SO\textsubscript{2} days (See Table S1 for a total list of strike days, and Figure 1). Several methods of addressing this issue were tested: no adjustment, excluding strike days, or adjusting for the indicator. The latter was found to yield models with a better fit and was used where appropriate. We used an autoregressive term (adjusting for the outcome at lag 1) to improve the autocorrelation of the model residuals. (32) The terms \( s_1(\text{Temperature}) \) and \( s_2(\text{day in the time series}) \) are smoothing functions which allow for non-linearity. The term \( s_3(\text{day of the year, } bs=\text{"cc"}) \) is a b-spline with a penalized cyclical cubic (a spline whose ends match up, and is thus suitable for modeling annual fluctuations) term day of year designed to control for seasonal trend. (33) Results from models with no adjustment for other pollutants (Partially adjusted models) are also presented. Quasipoisson distribution was assumed for all outcomes. All analysis was performed in R Studio. (34)

Subgroup analysis
We present results from stratified analyses of categories of anti-asthma drugs; 1) adrenergic inhalants (R03A), a large proportion of which are short-acting beta agonists, and 2) other inhalant drugs (R03B), mainly glucocorticoids and anticholinergenics. For PCMD and HED visits, we present results stratified for 1) infectious diseases including acute upper respiratory infections (World Health Organisation International Classification of Disease (ICD) codes J00-J06), influenza and pneumonia (J09-J18), and other acute lower respiratory infections (J20-J22), and 2) obstructive respiratory disease, including chronic obstructive pulmonary disease, COPD, and asthma (J44 and 45). Age-stratified results are also presented.

**Sensitivity analysis**

To explore the robustness of the analysis, we analysed the association between all respiratory PCC contacts including phone calls, consultations, and “other” because PCMD visits are subject to availability. To estimate the total impact on the health care systems from increased population morbidity we also analysed all PCC contacts including recurring PCC contacts within 14 days. For HED visits, we performed a sensitivity analysis including only individuals who were admitted for in-patient care (Table S3). Additionally, we performed sensitivity analysis on different lags of SO₂ for HED admissions, as our exploratory analysis revealed lag-specific effects for different age-categories (Figure S2c-d, Table S3). In sensitivity analysis of the exposure (Table S4), we wished to exclude the possibility of our results being entirely due to official advice encouraging individuals with respiratory diseases to have sufficient medicine at hand. This was broadcast to the public on days with forecasts of high SO₂ and it was speculated that some SO₂-associated increases in AMD could be due to compliance with this advice. As AMD is most often dispensed in larger quantities, a compliance effect would present itself as decreased effect estimates of SO₂ after the first days or week of warnings as regular users have filled their supplies. Thus, present results from analyses where we 1) excluded the first day, then 2) the first week, from the time series (Table S4). To eliminate any confounding effect of air pollution from other volcanic eruptions ((Eyjafjallajökull 2010 and Grímsvötn 2011) that impacted air quality during the study period (Daily mean of key air pollutants and events are indicated in Figure S1)(35) we reanalysed AMD, PCMD and HED excluding the years 2010 and 2011 (Table S4). Also, we performed analyses allowing for non-linearity of both lag structure, and concentration of SO₂ and present those results in the supplement (Figure S2 and Figure S3). Furthermore, we investigated lag-responses of sub-categories of respiratory disease (Figure S4) and effects of longer lags (Figure S5).

**Results**

The daily mean SO₂ concentrations in Iceland’s capital area were low or moderate until the Holuhraun eruption began 30 August 2014. After the eruption began, 24-hour SO₂ concentrations surpassed the air quality guideline limit of 125 µg/m³ on ten days. Both mean and median concentrations were significantly higher than before the eruption; mean 35.7 µg/m³ (SD 71.2) vs mean 1.4 µg/m³ (SD 1.1)
(Table 1, see also Figure 2 and Figure S1), and the correlations between air pollutants were altered substantially after the eruption (Table S1).

Before the eruption, an average of 129.4 individuals per day were registered purchasing (=were dispensed) AMD. There were 56.3% females and the mean age was 35.7 (SD 22.9) years. In primary care, the most common form of contact was GP visits (61%) followed by phone calls (33%), the daily mean number of individuals with an MD visits in primary care (PCMD) for respiratory disease was 142.2, the mean age was 32.8 (SD 22.5) years and 60.9% were women. In HED, the daily mean number of individual visits for respiratory health outcomes were 18.3, the mean age was 36.3 years (SD 29.4) and 49.8% were women. After the eruption began, 158.4 individuals per day were dispensed AMD, the mean age was 44.5 (SD 26.3) years. 144.8 individuals per day visited an MD in primary care for respiratory disease, the mean age was 35.0 years (SD 26.6), 60.4% were women. In HED, 17.5 individuals attended for respiratory disease, the individuals had a mean age of 39.4 years (SD 30.5) and 51.8% were women (Table 2).

Comparing the daily number of events, only mean daily number of individuals with AMD was significantly increased compared with the reference period (129.4 vs 158.4, \( p < 0.001 \)). Restricting this analysis to only the time of the year where the eruption was ongoing (using September to December in the years 2012 and 2013 where there were no volcanic eruptions) we found that AMD was significantly increased 138.2 vs 158.4, \( p = 0.0035 \)). Neither total nor age categories of HED and PCMD MD visits were significantly higher or lower during the eruption period as a whole compared to the period before (Table 2, Figure 2).

Regression analysis results

We found that for AMD, the overall time trend adjustment without fixed degrees of freedom showed signs of overfitting and so, comparing the model fits (\( R^2 \) and explained deviance), 4 degrees of freedom was selected as the optimal value for the time trend spline in analyses of AMD. For PCMD and HED, the degrees of freedom were not fixed. After inspecting the residuals for signs of drift, we were confident that the time trends adjust for the population increase during the study period.

In both partially (model 1) and fully adjusted (model 2) models, \( \text{SO}_2 \) was associated with increased number of AMD (model 2) by 0.99% (95% CI 0.39 – 1.58%) per 10 µg/m\(^3\) \( \text{SO}_2 \). High \( \text{SO}_2 \)-days, with concentrations exceeding the air quality guideline of 125µg/m\(^3\) were associated with a statistically significant increase in AMD by 10.9% (95% CI 2.1 – 19.6%) at lag 0–2 (Table 3). For PCMD and HED, the strike indicator for days where MDs were on strike was associated with fewer outcome events (as fewer MDs were working). The inclusion of the strike indicator variable improved the model fit and was an effect modifier of \( \text{SO}_2 \), increasing the effect estimates, and thus it was added to the following PCMD and HED models. In primary care, \( \text{SO}_2 \) was associated by an increase in PCMD visits by 1.26 % (95% CI 0.72 – 1.80%) per 10 µg/m\(^3\) \( \text{SO}_2 \) at lag 0–2. For PCMD, exposure to \( \text{SO}_2 \) over 125µg/m\(^3\) was associated with increases by 17.2 % (95% CI 10.0 – 24.4%) at lag 0-2. For HED visits, only adjusted results (model 2) of continuous \( \text{SO}_2 \) exposure at lag 0–2 yielded significant associations associated with increase (Figure
S2d) in total HED by 1.02% (95% CI 0.02 – 2.03%) per 10 µg/m³. There were no statistically significant associations between high SO₂ days and total HED (Table 3). The effect estimates from the partially adjusted models (model 1) were within the confidence intervals of the model 2 fully adjusted results (~10 percentage point change), except for HED visits which increased from 0.83% (95% CI -0.12 – 1.78%) in the unadjusted model 1 to 1.02% (95% CI 0.02 – 2.03%) per 10 µg/m³ SO₂ in model 2 (Table 3). Fully adjusted model results are presented from here.

In subgroup analysis (Table 4), the dispensing of short acting bronchodilator medication was associated with SO₂ exposure by 1.09% (95% CI 0.49-1.70%) per 10 µg/m³ SO₂, and 13.7% (95% CI 4.5-23.7%) after high SO₂ days. SO₂ exposure was not significantly associated with dispensing of long-acting corticosteroid medication. SO₂ exposure was associated with increased PCMD visits for respiratory infection diagnoses by 1.12% (95% CI 0.54 – 1.71%) and obstructive pulmonary disease by 2.08% (95% CI 1.13 – 3.04%) per 10 µg/m³ at lag 0–2, and PCMD visits for %). Following high SO₂ days, the estimated increase in PCMD visits were 15.8% (95% CI 7.2 – 25.1%) and 28.4% (95% CI 12.8 – 46.2%) for respiratory infections and obstructive disease, respectively. For HED visits, SO₂ exposure was not associated with significant increases in visits due to respiratory infections or asthma or COPD although all estimates were numerically indicative of a positive effect (Table 4, Figure S4).

**Age- specific effects**

In analysis stratified by age, the estimated association between SO₂ and AMD exposure at lag 0-2 were highest in children, and lower in elderly, however, all confidence intervals overlapped in both partially and fully adjusted models (Table 5). In PCMD, the effect estimates were highest in adults, and was not statistically significant in elderly.

**Outcome definitions of PCC**

Changing the outcome to include all primary care contacts (also phone calls) and further including the recurring contacts for the same diagnostic category within 14 days only marginally alter the results (17.2% vs 17.4% vs 20.3%) for high SO₂ levels (Table S3).

**Age-and lag-days**

Considering effects on HED of SO₂ exposure at lag 2-4, the effect estimated in children increased numerically from 1.22 % (95% CI -0.62 – 3.10%) to 2.21% (95% CI 0.30 – 3.97%) however, the confidence intervals overlapped and statistical power was low (Table S3).

Confining the analysis of HED to those who were admitted, there were no associations between SO₂ exposure at lag0-2, but admission in children were increased at lag 2-4 by 6.6% per 10 µg/m³ SO₂ (Table S3).

**Compliance with official advice, other eruptions**
Excluding the first day in a series of high SO\textsubscript{2} days from the exposure variable yielded very similar estimated effects of SO\textsubscript{2} exposure to the main analysis for AMD, but slightly lower for PCMD (Table S4). Excluding the whole week after the first high SO\textsubscript{2} concentration day, effect estimates for PCMD visits were higher in elderly compared with the main analysis (Table S4). Excluding the years 2010 and 2011 yielded statistically significant effect estimates for total AMD and PCMD, but only HED in elderly was significantly increased (Table S4). In the analysis of HED admissions rather than all HED visits, all effect estimates were positive, but there were no significant associations between SO\textsubscript{2} and HED at lag 0-2. When increasing the lag-period to 20 days we observed a second increase in PCMD at lag 16 after the initial SO\textsubscript{2} exposure, and at lag 11-13 for HED in young people and adults (Figure S4).

**Discussion**

Although SO\textsubscript{2} pollution levels were overall higher after the Holuhraun eruption began (Table 1), a crude comparison of respiratory health care utilisation in Iceland’s capital area before and after the beginning of the Holuhraun eruption (Table 2) showed that only AMD increased significantly compared to the corresponding months in previous years, as well as compared to the total control period. However, in time series regression analysis (Table 3), SO\textsubscript{2} concentrations were associated with increased AMD, PCMD and HED for respiratory causes at lag 0-2. Analysing the association with SO\textsubscript{2} levels which exceeded the 24-hour health limit (125 µg/m\textsuperscript{3}) to quantify the risk associated with high levels yielded similar results, and spline plots suggest a linear dose-response curve (Table 3, see also Figure S3). Stratifying AMD into subgroups, only dispensing of short-acting medications were significantly increased, indicating that dispensing of short-acting drugs are a more sensitive indicator of an immediate need for symptom relief, although the use of short-acting drugs are not specific to asthma disease (Table 4). In PCMD we found that PCMD for respiratory infections and obstructive lung disease were increased and the increase in obstructive lung disease suggested that individuals with this diagnosis were increasingly in need of care following SO\textsubscript{2} exposure (Table 4). For COPD diagnoses in HED, our results indicated a positive association, but it was lower than for PCMD, and the results did not reach statistical significance.

**Age-specific**

Stratifying by age-categories, the results indicated that for AMD, we observed with highest risk estimates with SO\textsubscript{2} exposure in individuals under 18, for PCMD, adults had highest risk, and for HED the highest risk was in individuals 65 and older (Table 5), except for lag2-4, where it was higher in individuals under 18 (Table S3). However, the confidence intervals overlapped, and the association between SO\textsubscript{2} and HED in elderly was not significant after adjusting for other pollutants than SO\textsubscript{2} (Table 5).

**Outcomes**

Exploring definitions of outcomes in primary care in sensitivity analyses, the tendency towards increasing effect estimates in elderly in analyses of all contacts and recurring contacts (Table S3) rather than non-recurring MD visits suggests that the care burden is higher in this group, but the confidence intervals
overlapped with those in the main analysis (Table 5). In HED admissions, a high effect estimate was seen for both continuous SO$_2$ and high SO$_2$ days in children at lag2-4, but the confidence intervals were wide (Table S3).

**Age-and lag specific**

In the regression analysis, the choice of lags were based on inspection of lag structures generated by lag-splines (Figure S2). For AMD and PCMD visits, the observed effects of SO$_2$ exposure occurred during the same day and up to two days after a peak in SO$_2$ concentration (Figures S2a-b). However, for HED visits in elderly, the observed effects occurred already at lag 0 (Figure S2c-d). A possible explanation for this is that primary care is the first point of contact for most individuals and hospital care is sought only after other care options have been exhausted, but for elderly, hospital care is sought immediately, possibly due to a more severe presentation of symptoms or complications due to underlying diseases, as illustrated by a significant association between SO$_2$ and HED in elderly before adjusting for other pollutants at lag0-2 (Table S3), and significant association only with HED in children at lag 2-4 (Table S3). Unfortunately, we did not have access to information about comorbidities and were unable to adjust for this.

**Compliance, other eruptions, longer lags**

In the sensitivity analysis to investigate the influence of compliance with official advice, the effect of initial warnings of volcanic gas exposure appeared to be limited, as excluding the first day or week of high SO$_2$ yielded similar estimates for AMD and PCC visits as the main analysis, although there was some loss of statistical power (Table S4). We speculate that since the increases in AMD and PCC persisted after the first exposed day and weeks, this indicates that the observed increases in health care utilisation reflect actual increased respiratory morbidities rather than merely adherence to official advice (Supplemental Table S4). Excluding the years 2010 and 2011 from the analysis to exclude effects of volcanic ash from Eyjafjallajökull and Grímsvötn yielded results within confidence intervals of our main analysis (Table S4). Investigating longer lags, we observed statistically significant increases in total PCMD and HED visits in young and adults at more than 15 days delay, but in both cases, these follow periods of decreases in outcomes, and are challenging to interpret (Figure S5).

**Relate to literature**

Although it has been suspected that the complex mixture of a volcanic plume (7) could have different health effects than merely SO$_2$, we found that our results were consistent with concentration-response functions found in studies from urban settings; e.g. respiratory mortality rates were estimated to increase by 2.4% per 27 µg/m$^3$ SO$_2$ (Li et al. 2017), corresponding to a point estimate of 0.88% per 10 µg/m$^3$ SO$_2$, lower, but within the confidence interval of the estimated increases of HED found in our study, namely 1.02% (95% CI 0.02 – 2.03%) per 10 µg/m$^3$. In previous studies of SO$_2$ exposure during volcanic
eruptions, the SO₂ concentration on Miyakejima Island increased after Mount Oyama erupted in 2000 meaning that residents and aid workers returning to the island from 2005 and onwards were exposed. Children living on the island who were exposed to daily mean concentration of 125 µg/m³ SO₂, had increased rates of wheezing.(36) In follow-up studies, permanent residents of Miyakejima Island (n=168) who lived in areas with high SO₂ exposure reported increased rates of cough and wheeze(13), both symptoms of asthma. None of the studies of people exposed at Miyakejima Island found adverse effects on lung function.(12, 14, 36)

Regarding individual susceptibility, we observed the highest effect estimates for PCMD visits for asthma and COPD (Table 4), indicating that individuals with these diseases are at increased risk, which is. Hyper-responsiveness to SO₂ has previously been reported as common (20-25%) in individuals with positive asthma test (methacholine test), indicating that they may be particularly vulnerable to severe SO₂ exposure.(4) Non-smokers have previously been reported as being at risk of symptoms after volcanic SO₂ exposure,(11) but unfortunately, smoking status was not available in our data.

Respiratory infections were increased in our study and similarly rates of cough and acute pharyngitis diagnosed at a clinic in SO₂-exposed communities near The Kilauea Volcano in Hawaii increased after a eruption activity increased in 2008.(18) Respiratory infection diagnoses in PCMD were statistically significantly associated with SO₂, a similar association has previously been reported for hospital visits for upper respiratory tract infections.(37) Volcanic eruptions have been associated with altered rates of other health outcomes such as accidents and mortality,(38) but these fell outside the scope of this study.

**Methodology**

Our study employs a time series design were individual level risk factors are not time-dependent and thus should not confound the association between short term exposure to air SO₂ and health outcomes, leaving bias due to unmeasured confounders, seasonal variation, and other intermittent exposures as main concerns. Previous studies of health effects of air pollution in Iceland have yielded lower effect estimates of daily air pollution on morbidity than the current study,(39, 40) other air pollution exposure types are thus not likely to bias the results. This includes H₂S, which has a low correlation (<0.1) with the exposure of interest during the study period (data not shown). During the reference period, there were two ash-rich volcanic eruptions, in Eyjafjallajökull 2010 and Grímsvötn 2011. While ash from Eyjafjallajökull had local respiratory health effects(24) and there may have been adverse health effects in the capital area,(40) neither eruption had significant SO₂ emissions and sensitivity analysis excluding this period yielded results within the confidence interval of the main study results (Supplemental Table S5). The exposed period October and November of 2014 did not coincide with any viral respiratory illnesses (e.g. influenza and RS-virus) epidemic,(41, 42) but it did coincide with the MD labour conflict which resulted in lower PCMD and HED attendance during those days. Hence, results from the analysis comparing the
eruption period with the time before may have underestimated of true effect of the SO$_2$ from the eruption.

**Strengths and limitations**

SO$_2$, PM$_{10}$ and NO$_2$ data was missing for a number of days which were excluded from the analysis. As the volcanic plume effectively changed the chemical composition of the atmosphere during the eruption period (Supplemental Tables S1a-b), the correlations of SO$_2$ with other air pollutants were altered after the eruption. Results that were adjusted for other pollutants were nearly identical to the main analysis (Table 3), indicating that the effect of the very high concentrations of SO$_2$ was unambiguous.

It is a strength of the current study that health data were collected prospectively from population-wide registers, which minimizes the risk of information bias from individuals knowing their exposure status. Although the exposure would have been known to the public for at least part of the exposed period but we attempt to address this source of bias in the unadjusted analysis and found only moderate changes to the results (Table S3). As a study outcome, we use dispensing of asthma-medication, a novel and more sensitive proxy for respiratory health in a population (28) than primary care attendance and hospital visits. It targets individuals who are already sensitive to poor air quality (26, 29) who in most cases have contact with the health care system. A limitation with register-data is that the data are not collected for research purposes and diagnoses given in the health care system could be biased and lead to overestimation as medical professionals assume respiratory outcomes to be more likely during the eruption. However, as the estimated effect are similar across all respiratory outcomes we conclude that this source of bias is not likely to explain our findings. By defining exposure status based on residential postcode we make several assumptions, firstly, that the whole area is equally exposed, and secondly, that the study participants are physically near their homes. However, these sources of bias would result in wider confidence intervals or bias the results toward the null. Reykjavík and the Icelandic capital area was exposed to SO$_2$ concentrations above 125µg/m$^3$ during a total of ten days which occurred mostly during October and November 2014. This limits the statistical power of the study and our options for further analysis, as does the fact that our study period did not extend until after the eruption, meaning that we cannot fully assess the complete impact of the health impact from the eruption on the population from our results.

**Conclusion**

In conclusion, this comprehensive study with prospectively collected data on volcanic SO$_2$ air pollution exposure and respiratory health outcomes, is the first to firmly establish an association between spikes of high SO$_2$ concentrations and respiratory outcomes in the general population, and specifically in individuals with prevalent respiratory disease. These findings emphasize the need for attention from authorities and susceptible individuals during times of volcanic eruptions.
Abbreviations

AMD  Asthma medication dispensing
COPD  Chronic obstructive pulmonary disease
HED  Hospital emergency department
PCC  Primary care contacts
PCMD Primary care MD visits
SO₂  Sulfur dioxide
PM₁₀  Particle matter less than 10 µg/m³

Declarations

Ethics approval:

The study was approved by the Science Bioethics Committee of Iceland (reference: VSNb2015050022/03.01).

Data sharing statement

The research data contain sensitive individual-level information which is not publicly available. It can be made available to researchers after approval of a formal application to the Icelandic Directorate of Health and the Icelandic Bioethics committee. The exposure data is publicly available, please see http://umhverfisstofnun.is.

Acknowledgements

The authors wish to acknowledge The Environment Agency of Iceland and Kristinn Jónsson from The Icelandic Directorate of health for providing data and Dr. Evgenia Ilyinskaya for valuable comments on the manuscript. The study was funded by the Icelandic Ministry of Health. The funding source had no influence on the reporting or publication of the study.

Competing interests

The authors declare they have no actual or potential competing financial interests.

Funding

The study was funded by the Icelandic Ministry of Health.

Author contributions
TGu, UAV, HB, TGi conceived the study and contributed to the study design and manuscript. TJ and RGF provided exposure data, and contributed to the exposure assignment, RGF also contributed reporting of the methods. FD and TA contributed to the statistical analysis. HKC managed the study data, analysed the data, wrote the first draft of the manuscript and managed the revisions. All authors contributed to the manuscript.

Acknowledgements

The authors wish to acknowledge the The Environment Agency of Iceland and Kristinn Jónsson from the Icelandic Directorate of health for providing data, and Dr. Evgenia Ilyinskaya for valuable comments on the manuscript. The study was funded by the Icelandic Ministry of Health. The funding source had no influence on the reporting or publication of the study.

References

1. EPA. Integrated Science Assessment (ISA) for Sulfur Oxides—Health Criteria. Rep EPA/600/R-08/047F. 2008.
2. World Health Organization. WHO Air quality guidelines for particulate matter, ozone, nitrogen dioxide and sulfur dioxide: global update 2005: summary of risk assessment. World Health Organization; 2006.
3. Reiss R, Anderson EL, Cross CE, Hidy G, Hoel D, McClellan R, et al. Evidence of health impacts of sulfate-and nitrate-containing particles in ambient air. Inhalation toxicology. 2007;19(5):419-49.
4. Nowak D, Jorres R, Berger J, Claussen M, Magnussen H. Airway responsiveness to sulfur dioxide in an adult population sample. American journal of respiratory and critical care medicine. 1997;156(4):1151-6.
5. Small C, Naumann T. The global distribution of human population and recent volcanism. Global Environmental Change Part B: Environmental Hazards. 2001;3(3):93-109.
6. Newnham RM, Dirks KN, Samaranayake D. An investigation into long-distance health impacts of the 1996 eruption of Mt Ruapehu, New Zealand. Atmospheric Environment. 2010;44(12):1568-78.
7. Ilyinskaya E, Schmidt A, Mather TA, Pope FD, Witham C, Baxter P, et al. Understanding the environmental impacts of large fissure eruptions: Aerosol and gas emissions from the 2014–2015 Holuhraun eruption (Iceland). Earth and Planetary Science Letters. 2017;472:309-22.
8. Gísladóttir, G. and G. Jóhannesdóttir (2016). Residentý risk perception of and response to SO2 risk in east Iceland during the volcanic eruption in Bárðarbunga/Holuhraun 2014-2015. EGU General Assembly. Vienna, Austria: EPSC2016-8137.
9. Gíslason SR, Stefansdottir G, Pfeffer M, Barsotti S, Jóhannsson T, Galeczka IM, et al. Environmental pressure from the 2014–15 eruption of Bárðarbunga volcano, Iceland. 2015.
10. Carlsen HK, Aspelund T, Briem H, Gislason T, Jóhannsson T, Valdimarsdóttir U, et al. Respiratory health among professionals exposed to extreme SO 2 levels from a volcanic eruption. Scandinavian
journal of work, environment & health. 2019;45(3).

11. Ishigami A, Kikuchi Y, Iwasawa S, Nishiwaki Y, Takebayashi T, Tanaka S, et al. Volcanic sulfur dioxide and acute respiratory symptoms on Miyakejima island. Occupational and Environmental Medicine. 2008;65(10):701-7.

12. Iwasawa S, Kikuchi Y, Nishiwaki Y, Nakano M, Michikawa T, Tsuboi T, et al. Effects of SO2 on respiratory system of adult Miyakejima resident 2 years after returning to the island. J Occup Health. 2009;51(1):38-47.

13. Iwasawa S, Nakano M, Tsuboi T, Kochi T, Tanaka S, Katsunuma T, et al. Effects of sulfur dioxide on the respiratory system of Miyakejima child residents 6 years after returning to the island. International archives of occupational and environmental health. 2015;88(8):1111-8.

14. Kochi T, Iwasawa S, Nakano M, Tsuboi T, Tanaka S, Kitamura H, et al. Influence of sulfur dioxide on the respiratory system of Miyakejima adult residents 6 years after returning to the island. J Occup Health. 2017:16-0256-OA.

15. Longo B, Rossignol A, Green J. Cardiorespiratory health effects associated with sulphurous volcanic air pollution. Public health. 2008;122(8):809-20.

16. Longo BM, Yang W. Acute bronchitis and volcanic air pollution: a community-based cohort study at Kilauea Volcano, Hawai’i, USA. J Toxicol Environ Health Part A. 2008;71(24):1565-71.

17. Longo BM. Adverse health effects associated with increased activity at Kilauea Volcano: A repeated population-based survey. International Scholarly Research Notices. 2013;2013.

18. Longo BM, Yang W, Green JB, Crosby FL, Crosby VL. Acute health effects associated with exposure to volcanic air pollution (vog) from increased activity at Kilauea Volcano in 2008. J Toxicol Environ Health Part A. 2010;73(20):1370-81.

19. The International Volcanic Health Hazard Network - IVHHN. Sulphur Dioxide (SO2). 2019. https://www.ivhhn.org/information/information-different-volcanic-gases/sulphur-dioxide/ Accessed 13 December 2019.

20. Mnatzaganian C, Lozano A, Pellegrin K, Miyamura J, Knox M, Hanlon A, editors. Acute Respiratory and Cardiovascular Outcomes Associated with Low Levels of Ambient Fine Particulate Matter (PM2.5) on the Island of Oahu. Proceedings of the 52nd Hawaii International Conference on System Sciences; 2019.

21. Hansell A, Oppenheimer C. Health hazards from volcanic gases: a systematic literature review. Archives of Environmental Health: An International Journal. 2004;59(12):628-39.

22. Michellier C, Katoto PdMC, Dramaix M, Nemery B, Kervyn F. Respiratory health and eruptions of the Nyiragongo and Nyamulagira volcanoes in the Democratic Republic of Congo: a time-series analysis. Environmental Health. 2020;19(1):1-11.

23. Iceland S. Population by postal code, sex and age 1998-2017 [Mannfjöldinn eftir póstnúmerum, kyni og aldri 1998-2017]. 2017.
24. Sigurgeirsdóttir S, Waagfjörð J, Maresso A. Iceland: Health system review. *Health systems in Transition*. 2014;116(6):1-182.

25. Carlsen HK, Hauksdottir A, Valdimarsdottir UA, Gíslason T, Einarsdottir G, Runolfsson H, et al. Health effects following the Eyjafjallajökull volcanic eruption: a cohort study. BMJ open. 2012;2(6).

26. Menichini F, Mudu P. Drug consumption and air pollution: an overview. *Pharmacoepidemiology and drug safety*. 2010;19(12):1300-15.

27. Furu K, Skurtveit S, Langhammer A, Nafstad P. Use of anti-asthmatic medications as a proxy for prevalence of asthma in children and adolescents in Norway: a nationwide prescription database analysis. *European journal of clinical pharmacology*. 2007;63(7):693-8.

28. Naureckas ET, Dukic V, Bao X, Rathouz P. Short-acting β-agonist prescription fills as a marker for asthma morbidity. *Chest*. 2005;128(2):602-8.

29. Fattore E, Davoli E, Castiglioni S, Bosetti C, Depaolini AR, Marzona I, et al. Wastewater-based epidemiological evaluation of the effect of air pollution on short-acting beta-agonist consumption for acute asthma treatment. *Environmental research*. 2016;150:106-11.

30. Gasparrini A, Armstrong B, Kenward MG. Distributed lag non-linear models. *Statistics in medicine*. 2010;29(21):2224-34.

31. Dominici F, Peng RD. Statistical methods for environmental epidemiology with R: a case study in air pollution and health: *Springer*; 2008.

32. Brumback BA, Ryan LM, Schwartz JD, Neas LM, Stark PC, Burge HA. Transitional regression models, with application to environmental time series. *Journal of the American Statistical Association*. 2000;95(449):16-27.

33. Bhaskaran K, Gasparrini A, Hajat S, Smeeth L, Armstrong B. Time series regression studies in environmental epidemiology. *International journal of epidemiology*. 2013;42(4):1187-95.

34. R Core Team R. R: A language and environment for statistical computing. *R foundation for statistical computing* Vienna, Austria; 2013.

35. Thorsteinsson T, Jóhannsson T, Stohl A, Kristiansen NL. High levels of particulate matter in Iceland due to direct ash emissions by the Eyjafjallajökull eruption and resuspension of deposited ash. *Journal of Geophysical Research: Solid Earth*. 2012;117(B9).

36. Iwasawa S, Michikawa T, Nakano M, Nishiwaki Y, Tsuboi T, Tanaka S, et al. Nine-month observation of effects of SO₂ on the respiratory system in child Miyakejima citizens. [Nihon koshu eisei zasshi] *Japanese journal of public health*. 2010;57(1):39-43.

37. Li R, Jiang N, Liu Q, Huang J, Guo X, Liu F, et al. Impact of air pollutants on outpatient visits for acute respiratory outcomes. *International journal of environmental research and public health*. 2017;14(1):47.

38. Fano, V., Cernigliaro, A., Scondotto, S., Perucci, C. A., & Forastiere, F. (2010). The fear of volcano: short-term health effects after Mount Etna’s eruption in 2002. *European Respiratory Journal*, 2010;36(5), 1216-1218.
Tables

Table 1 Air pollution in Iceland’s capital area before and during the Holuhraun eruption

| Pollutant | Before eruption 2010-01-01 - 2014-08-30 1703 days | During eruption 2014-08-31 - 2014-12-31 123 days |
|-----------|-------------------------------------------------|--------------------------------------------------|
| SO₂       | Mean (SD) 1.4 (1.1) | Range 0.0-20.6 | Days with missing data 20 | Mean (SD) 35.7 (71.2) | Range 1.3-418.0 | Days with missing data 0 | p* <0.001 |
| PM₁₀      | Mean (SD) 20.3 (18.2) | Range 3.8-315.4 | Days with missing data 19 | Mean (SD) 17.8 (9.1) | Range 5.7-71.5 | Days with missing data 13 | p* 0.010 |
| NO₂       | Mean (SD) 15.8 (11.3) | Range 0.0-75.9 | Days with missing data 54 | Mean (SD) 12.0 (8.2) | Range 0.4-41.0 | Days with missing data 0 | p* <0.001 |

*from a t-test of means. SD: Standard deviation.

Table 2 Demographic characteristics of the study population and daily number of individuals who were dispensed anti-asthma medication (AMD) or utilized health services for respiratory disease before and during the Holuhraun eruption
| Anti-asthma medication dispensions (AMD)** | Before eruption | During eruption |
| --- | --- | --- |
| | n | % | (mean) | Range | n | % | (mean) | Range | P* |
| All | 45 | 100% | 129.4 | 5-715 | 11 | 100% | 158.4 | 22-301 | <0.001 |
| Female sex | 25 | 56.3% | 75.1 | 2-430 | 6 | 59.1% | 93.2 | 13-185 | <0.001 |
| Under 18 years of age | 14 | 31.1% | 28.3 | 0-139 | 2 | 21.8% | 33.2 | 8-68 | <0.001 |
| 18-64 years of age | 24 | 53.0% | 60.7 | 3-386 | 5 | 59.7% | 75.2 | 5-149 | <0.001 |
| 65 years and older | 7 | 15.9% | 40.4 | 0-202 | 3 | 27.5% | 50.1 | 0-105 | <0.001 |

| Primary care center MD visits (PCMD) | All | Female sex | Under 18 years of age | 18-64 years of age | 65 years and older |
| --- | --- | --- | --- | --- | --- |
| | n | % | (mean) | Range | n | % | (mean) | Range | P* |
| All | 106 | 100% | 142.2 | 17-325 | 15 | 100% | 144.8 | 41-259 | 0.6531 |
| Female sex | 59 | 55.7% | 84.6 | 9-180 | 9 | 60.4% | 87.9 | 22-165 | 0.343 |
| Under 18 years of age | 31 | 29.3% | 39.2 | 3-136 | 4 | 26.2% | 38.0 | 13-75 | 0.4019 |
| 18-64 years of age | 64 | 60.9% | 85.9 | 12-191 | 9839 | 62.2% | 89.7 | 24-161 | 0.2951 |
| 65 years and older | 10 | 9.8% | 17.1 | 0-48 | 1 | 11.6% | 17.1 | 0-42 | 0.9877 |

| Hospital emergency department visits (HED) | All | Female sex | Under 18 years of age | 18-64 years of age | 65 years and older |
| --- | --- | --- | --- | --- | --- |
| | n | % | (mean) | Range | n | % | (mean) | Range | P* |
| All | 19 | 100% | 18.3 | 1-45 | 1 | 100% | 17.5 | 4-34 | 0.176 |
| Female sex | 9 | 49.8% | 9.2 | 0-29 | 1000 | 51.8% | 9.0 | 2-18 | 0.527 |
| Under 18 years of age | 6 | 34.9% | 5.7 | 0-24 | 638 | 33.0% | 9.0 | 0-16 | 0.640 |
| 18-64 years of age | 8 | 42.3% | 7.0 | 0-24 | 758 | 39.2% | 5.5 | 1-14 | 0.269 |
| 65 years and older | 4 | 22.9% | 5.7 | 0-19 | 536 | 27.7% | 6.7 | 0-13 | 0.198 |

*from a t-test of means.

Table 3 Percent excess risk associated with daily SO$_2$ exposure at lag -2 (for exposure as a continuous variable, left, and as an indicator
or days with pollution levels above the air quality guideline value, and respiratory health outcomes in the capital area of Iceland

| SO$_2$ (per 10 µg/m$^3$) | SO$_2$ levels $>$125 ug/m$^3$ |
|--------------------------|-----------------------------|
| % | 95% CI | % | 95% CI |
| **Asthma medication (AMD)** | | | |
| Model 1 | 0.89% | 0.29% | 1.49% | 10.5% | 1.8% | 20.0% |
| Model 2 | 0.99% | 0.39% | 1.58% | 10.9% | 2.1% | 19.6% |
| **Primary care MD visits (PCMD)** | | | |
| Model 1 | 1.18% | 0.68% | 1.69% | 16.9% | 9.2% | 25.0% |
| Model 2 | 1.26% | 0.72% | 1.80% | 17.2% | 10.0% | 24.4% |
| **Hospital Emergency Department (HED)** | | | |
| Model 1 | 0.83% | -0.12% | 1.78% | 11.7% | -1.4% | 26.7% |
| Model 2 | **1.02%** | **0.02%** | **2.03%** | 9.4% | -3.8% | 22.6% |

Model 1: Adjusted for season, time trend, day of week, odd holidays, the outcome at lag 1, temperature, and relative humidity. n=1606

Model 2: Adjusted for season, time trend, day of week, odd holidays, the outcome at lag 1, temperature, relative humidity, NO$_2$, PM$_{10}$ at the same lags as SO$_2$. n=1529.

Table 4 Percent excess risk associated with SO$_2$ exposure at lag 0-2 both for exposure as a continuous variable and for days with pollution levels above the air quality guideline value) and changes in respiratory health in diagnosis subcategories in primary care and hospital emergency departments in the capital area of Iceland.
|                                | Mean (SD) | SO₂ (per 10 µg/m³)* | SO₂ levels >125 SO₂ µg/m³ | 95% CI | 95% CI |
|--------------------------------|-----------|---------------------|-----------------------------|--------|--------|
| **Anti-asthma medication**     |           |                     |                             |        |        |
| Short-acting β-agonist (R03A)  | 88 (47.2) | 1.09% 0.49% 1.70%   | 13.7% 4.5% 23.7%            |        |        |
| Long-acting (R03B)             | 38.7 (21.4)| 0.74% -0.03% 1.52% | 8.7% -2.2% 20.7%            |        |        |
| **Primary care MD visits (PCMD)** |          |                     |                             |        |        |
| Respiratory infections (J0-J2) | 108.6 (46.2)| 1.12 0.54% 1.71%   | 15.8% 7.2% 25.1%            |        |        |
| Asthma and COPD (J44-45)       | 15.6 (9.6)| 2.08% 1.13% 3.04%   | 28.4% 12.8% 46.2%           |        |        |
| **Hospital emergency department visits (HED)** | |                     |                             |        |        |
| Respiratory infections (J0-J2) | 9.3 (4.3) | 1.09% -0.22% 2.41% | 7.9% -60.9% 27.8%           |        |        |
| Asthma and COPD (J44-45)       | 4.3 (2.7) | 0.95% -0.99% 2.93% | 3.3% -19.8% 33.2%           |        |        |

* Adjusted for season (spline), time trend (spline), day of week, odd holidays 1, temperature (spline), relative humidity, NO₂, PM₁₀ at the same lags as SO₂, and the outcome at lag.

Table 5 Percent excess risk in age-specific respiratory health outcomes associated with SO₂ exposure at lag 0-2 (both for exposure as a continuous variable and for days with pollution levels above the air quality guideline value).
| Anti-asthma medication (AMD) | SO₂ (per 10 µg/m³) | 95% CI | SO₂ levels >125 µg/m³ | 95% CI |
|-----------------------------|---------------------|--------|-----------------------|--------|
|                              | %                   |        | %                     |        |
| Children 0-17                | 1.48%               | 0.61%  | 2.35%                 | 20.2%  | 7.8%  | 32.6% |
| Adults 18-65                 | 0.86%               | 0.20%  | 1.53%                 | 11.3%  | 1.8%  | 20.9% |
| Elderly >65                  | 0.90%               | 0.14%  | 1.67%                 | 9.8%   | -1.4% | 20.9% |
| Primary care MD visits (PCMD) |                     |        |                       |        |
| Children 0-17                | 1.24%               | 0.40%  | 2.08%                 | 14.2%  | 3.0%  | 25.5% |
| Adults 18-65                 | 1.21%               | 0.63%  | 1.79%                 | 18.5%  | 10.6% | 26.3% |
| Elderly >65                  | 1.11%               | 0.14%  | 2.09%                 | 11.8%  | -2.0% | 25.6% |
| Hospital Emergency Department (HED) |                 |        |                       |        |
| Children 0-17                | 1.22%               | -0.62% | 3.10%                 | 7.0%   | -18.7% | 32.7% |
| Adults 18-65                 | 0.52%               | -0.99% | 2.06%                 | 0.0%   | -20.7% | 20.7% |
| Elderly >65                  | 1.01%               | -0.62% | 2.66%                 | 19.7%  | -1.8% | 41.2% |

All results are adjusted for season, time, day of week, odd holidays, the outcome at lag1, and MD strike for PCC and HED, relative humidity, temperature, PM₁₀, and NO₂. n=1788.

**Figures**
Figure 1

For each outcome, we constructed daily time series starting 1 January 2010 to 31 December 2014 for the following age groups; children (under 18 years of age), adults (18-64 years), and elderly (age 65 years and above), see data selection in Flow diagrams 1-3 in the supplemental material. We obtained SO2, PM10, and NO2 data along with meteorological data from the Icelandic Environment Agency’s stationary air pollution monitor located in Reykjavik.
Comparing the daily number of events, only mean daily number of individuals with AMD was significantly increased compared with the reference period (129.4 vs 158.4, p<0.001). Neither total nor age categories of HED and PCMD MD visits were significantly higher or lower during the eruption period as a whole compared to the period before (Table 2, Figure 2).

**Figure 2**

Supplementary Files
This is a list of supplementary files associated with this preprint. Click to download.

- SupplementalmaterialHoluhraunstudy201031.docx