Research article

Characterization of airborne dust samples collected from core areas of Kathmandu Valley

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\textbf{ABSTRACT}

Kathmandu Valley is reported to be one of the highly polluted and populated cities in the world. Particulate matter is one of the major contributors of unhealthy air in Kathmandu. Although there are several reports on spatial and temporal variation of air quality of Kathmandu Valley, the morphological and mineralogical characteristics of particulate matter are very limited or none. In this study, we report on the mineralogical and morphological analysis of airborne particulate matter collected from densely populated core areas of Kathmandu Valley using spectroscopic and microscopic techniques. The Fourier Transform Infrared (FTIR) and X-ray Diffraction (XRD) data showed the presence of clay minerals, crystalline silicate mineral, carbonate minerals, and asbestiform mineral in the dust samples. The field emission scanning electron microscopic analysis confirmed the existence of particles having diverse morphology with some of the particles having aspect ratio as high as twenty; indicating the existence of asbestiform type minerals. Based on SEM-EDX data, we found that the relative distribution of elements to be different in different samples and C, O, Mg, Ca, and Si were the major elements in the dust samples. Interestingly, the XRD data analysis showed that in all the samples quartz mineral having high degree of crystallinity was present. The XRD measurement was also carried out in three different brands of cement samples. Few minerals present in dust samples were also identified in the cement samples. This observation could indicate that cement is one of the sources of minerals in the airborne particulate matter in the Kathmandu Valley.

1. Introduction

Air pollution is one of the major threats to climate, human health, and ecosystems and is one of the highly discussed global issues \cite{1, 2, 3, 4}. It is reported that the combined effect of outdoor and indoor air pollution accounts for around 7 million global deaths annually \cite{5}. According to the most recent air quality database, around 97\% of the cities in the low- and middle-income countries and 49\% of cities in high income countries having population more than 100,000 do not meet the world health organization (WHO) air quality guideline \cite{5}.

The airborne particles of size 10 $\mu$m or less (PM$_{10}$) and 2.5 $\mu$m or less (PM$_{2.5}$) are one of the major air pollutants that can penetrate deeply into the respiratory system. Exposure to these particles is linked to different health issues, such as chronic bronchitis, asthma, diabetes, stroke, cancer and eventually death \cite{5, 6, 7}. The global burden of disease (GBD) methodology estimated that ambient PM$_{2.5}$ pollution is the fifth leading cause of death causing 4.2 million deaths and 103.1 million disability adjusted life years in 2015 \cite{8}.

Health effect of particulate matter depends on multiple parameters such as exposure dose, chemical composition, morphological characteristics, surface reactivity, hydrophobicity and hydrophilicity, and solubility after deposition in the targeted sites \cite{9, 10, 11, 12, 13}. Inhalation of asbestiform particulate matter, depending on the nature and exposure dose, is reported to cause pleural fibrosis, asbestosis, carcinoma of lung, and mesothelioma \cite{10}. Similarly, inhalation of crystalline silica (quartz) is reported to tuberculosis, silicosis, chronic bronchitis/chronic obstructive pulmonary disease (COPD) and lung cancer while amorphous silica is found to be very less toxic \cite{14, 15}. Airborne particulate matter also contains clay minerals of different types. The inhalation of clay mineral rich particulate matter in occupational place is, however, reported to have minimal or no toxic effects \cite{16}. Airborne particulate matter having high content of heavy metals, such as lead, arsenic, and mercury is reported to cause health effects ranging from neurotoxicity to
In this research, we made mineralogical and morphological analysis of the airborne dust samples collected from eleven densely populated locations of Kathmandu Valley. We used FTIR and XRD measurement to identify the minerals. The information on the morphology and elemental composition of sample was obtained from SEM and SEM-EDX, respectively. We also analyzed XRD data of all the samples to identify the minerals. The information on the morphology and elemental composition of sample was obtained from SEM and SEM-EDX, respectively. We also analyzed XRD data of all the samples to identify the minerals.

2. Materials and methods

2.1. Sample collection

The passive sampling method was adopted in this study. The free-falling particles were collected in a sterile polystyrene petri dish (100 mm internal diameter and 15 mm depth, Fisher Scientific) in dry sunny days in March 2018. The petri dishes were placed at a height of 10 m above the ground in an open box. The samples were collected from eleven densely populated core locations of Kathmandu Valley. The samples were collected over a period of 24 h at each location. The sample collection locations are shown in Figure 1. The detail information on the sampling sites is provided in Table 1. The passive sampling method was adopted in this study. The free-falling particles were collected in a sterile polystyrene petri dish (100 mm internal diameter and 15 mm depth, Fisher Scientific) in dry sunny days in March 2018. The petri dishes were placed at a height of 10 m above the ground in an open box. The samples were collected from eleven densely populated core locations of Kathmandu Valley. The samples were collected over a period of 24 h at each location. The sample collection locations are shown in Figure 1. The detail information on the sampling sites is provided in Table 1.

2.2. Fourier transform infrared study

The infrared spectra of all the samples were collected in the range of 4000–400 cm⁻¹ by a Fourier transform infrared spectrometer (ABB Bomen MB100, Canada) in an attenuated total reflection mode (ATR). Before measuring the FTIR spectra of sample, background spectrum of ATR crystal was measured and subtracted from the sample spectrum. The spectral resolution of the spectrometer was 4 cm⁻¹. Each reported spectrum is the average of sixteen scans that provided excellent signal to noise ratio. The different minerals in the dust sample was identified on the basis of literature studies and RRUFF database of reference standard [37].

2.3. Electron microscopic study

The electron microscopic images and energy dispersive X-ray (EDX) spectra were measured by a field emission scanning electron microscope (FE-SEM, Carl Zeiss, Supra 40VP, Japan). For SEM measurement, the sample was mounted on aluminum stub using double sided carbon tape.

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Table 1. Details on sampling sites.

| Sample ID | Location name          | Location coordinates       |
|-----------|------------------------|---------------------------|
| $S_1$     | Bagbazar               | 27°42’22.93”N, 85°19’3.93”E |
| $S_2$     | Old Baneshwor          | 27°41’51.29”N, 85°20’3.57”E |
| $S_3$     | Ratna Rajya School, Baneshwor | 27°41’44.55”N, 85°20’14.74”E |
| $S_4$     | Kalanki                | 27°41’35.49”N, 85°16’57.58”E |
| $S_5$     | Putalisadak            | 27°42’20.52”N, 85°19’22.73”E |
| $S_6$     | Trichandra Campus, Ghantaghar | 27°42’28.21”N, 85°18’59.46”E |
| $S_7$     | Kalimati               | 27°41’57.57”N, 85°17’58.59”E |
| $S_8$     | Tripureshwor           | 27°41’38.81”N, 85°18’49.12”E |
| $S_9$     | Balaju                 | 27°43’38.64”N, 85°18’16.40”E |
| $S_{10}$  | Jawalakhel            | 27°40’22.18”N, 85°18’49.62”E |
| $S_{11}$  | Satdobato              | 27°39’32.48”N, 85°19’28.19”E |

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Figure 1. A part of Google Earth map showing sampling sites S1 to S11.
To make the sample surface conductive, a very thin layer of platinum was coated on the sample using a sputter coater. The sample was then placed in the vacuum chamber of SEM instrument and image was collected. For each sample 5–10 SEM images were taken at different magnifications. Image analysis was done in ImageJ software; a Java-based open source developed by National Institutes of Health and the Laboratory, USA. Aspect ratio of a particle was calculated by dividing the highest dimension (length) of the particle by its lowest dimension (width).

The SEM-EDX consisted of a silicon drift detector having detection limit of around 0.1% and capable of collecting spectrum from specific points and field of view for qualitative and semi-quantitative elemental analysis. In EDX measurement, we measured full field of view EDX spectra of all the samples. We also measured single particle spectra in few samples. The SEM image and the single particle EDX spectra for sample S2 and S3 are shown in Figure 3.

The atomic weight percentage distribution obtained from the EDX spectrum of all the samples is provided in Table 2.

2.3. FTIR measurement in the dust samples

We measured FTIR spectra of all the samples (S1 to S11) in attenuated total reflectance (ATR) mode. The FTIR spectrum of sample S11 is shown in Figure 4. For clarity, the spectrum collected in the middle IR (MIR) range is plotted in two separate frames, 400-1999 cm⁻¹ in frame A and 2000-4000 cm⁻¹ in frame B. To get more accurate peak/shoulder position in the range of 830–1250 cm⁻¹, a five component Gaussian fitting was used and the result is shown as inset in frame A.

The FTIR spectra of the remaining samples (S1–S10) in the range 400–4000 cm⁻¹ is provided in Figure 5.

2.4. X-ray diffraction measurement and analysis

The X-ray diffraction data were collected at Bragg angle 2θ ranging from 10 to 80° (step size 0.01°) by X-ray diffractometer (Rigaku, UK). The Cu Kα line having wavelength of 1.540 Å (30 kV, 40 mA) was used as X-ray source. The inter-planar spacing (d_hkl) was calculated by using the Bragg’s equation nλ = 2d_hklSinθ, where n is order of reflection (n = 1) and λ is wavelength of X-ray used (λ = 1.54Å). The parameters such as peak positions (2θ) and inter-planer spacing (d_hkl) were compared with the literature data and RRUFF database of reference standard to identify the different minerals present in the samples [37].

The crystallinity index (CI) of the quartz mineral in the samples was measured in the scale of 0–10 by using the formula CI = 10 aF/b, where F is experimental parameter (which is ~1.65), b is total intensity of peak at 67.8° (including amorphous background), and a is intensity of crystalline peak only [38].

3. Results

3.1. SEM and SEM-EDX measurement in the dust samples

We measured the FE-SEM images of all the samples (S1–S11) at different magnification. Representative images for the samples S1, S2, S3, S4, S9, and S10 are shown in Figure 2A and B.

We measured EDX data in samples S1–S11 and also the single particle EDX in the few selective particles/regions of the samples S2 and S3 to understand the elemental composition of the dust samples. The SEM image and the single particle EDX spectra for sample S2 and S3 are shown in Figure 3.

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3.2. FTIR measurement in the dust samples

We measured FTIR spectra of all the samples (S1 to S11) in attenuated total reflectance (ATR) mode. The FTIR spectrum of sample S11 is shown in Figure 4. For clarity, the spectrum collected in the middle IR (MIR) range is plotted in two separate frames, 400-1999 cm⁻¹ in frame A and 2000-4000 cm⁻¹ in frame B. To get more accurate peak/shoulder position in the range of 830–1250 cm⁻¹, a five component Gaussian fitting was used and the result is shown as inset in frame A.

The FTIR spectra of the remaining samples (S1–S10) in the range 400–4000 cm⁻¹ is provided in Figure 5.

3.3. XRD measurement

We measured the X-ray diffraction data of the dust samples and found that the XRD data for all samples were similar. The XRD datum of S11 is shown in Figure 6. For clarity, a separate plot for 20 value in the range 10–46.5° and 46.5–78° is made in frames A and B, respectively. For easy comparison, intensity is normalized at the most intense peak (2θ = 26.6°) and the peak height is truncated.

Minerals in the dust samples were identified by comparing the characteristic X-ray diffraction peaks to the literature data and an open source RRUFF database of reference standards [37]. We refer to section
For details on the assignment XRD peak and identification of minerals.

In Figure 6 the letters D is for mineral dolomite, C for calcite, Q for quartz, K for kaolinite, I for illite, H for hematite, V for vaterite, and Ch for chrysotile.

The XRD data of the samples S1–S10 is shown in Figure 7.

4. Discussion

4.1. FE-SEM and EDX measurements

The FE-SEM images indicated that most of the dust particles were symmetrical (lower aspect ratio in the range of 1–3) (Figure 2A) with few particles with diverse morphology (Figure 2B) and aspect ratio greater than three. For example, sample S3 consisted of sheet like particles having multiple micro fibrils and sample S9 ribbon shaped particles having aspect ratio as high as twenty. Such particles can be classified as asbestiform particles. This observation is in consistent to the recent study where particles of diverse morphology having aspect ratio as high as 20 was reported in dust samples of Kathmandu Valley [36].

On the basis of data in Table 2, we found that the dust samples contain metallic elements Fe, Mg, Al, Si, Ca, and Na, and non-metallic elements C and O distributed in different ratio. Calcium and silicon were the most abundant elements.

4.2. FTIR measurement in the dust samples

The Fourier transform infrared spectroscopy (FTIR) is one of the highly used techniques to characterize minerals present in dust samples [33, 39, 40, 41, 42]. The Si–O bending vibration modes at 465 cm$^{-1}$ and 694 cm$^{-1}$ and stretching modes at 777 cm$^{-1}$, 800 cm$^{-1}$, 1090, 1144
and 1872 cm$^{-1}$ are the characteristic frequencies of quartz [33, 40, 41, 42, 43, 44, 45]. These frequencies were observed in all the samples reported in this study (Figures 4 and 5). Quartz is one of the most abundant crystalline silicate minerals that contains a continuous framework of SiO$_4$ tetrahedra. The epidemiological studies have reported that inhalation of crystalline silica dust, depending on the exposure dose, can lead to inflammation, pulmonary tuberculosis, silicosis, and lung cancer [31, 46].

Clay minerals such as montmorillonite, kaolinite, and illite were reported in dust samples collected from different cities of the world. The absorption peaks near 3620 cm$^{-1}$ and 3550 cm$^{-1}$ were assigned to stretching frequency of O–H bond of H$_2$O bound directly to interlayer cations (inner sphere) and O–H of surface bonded water in montmorillonite [39]. The characteristic bending mode of AlAlO–H group in montmorillonite was reported to exist at 914 cm$^{-1}$. These characteristic frequencies were found in S1, S2, S3, S4, S5, S6, and S11 (Figures 4 and 5), thus indicating the presence of montmorillonite in these samples. Montmorillonite is a pylllosilicate group of clay minerals (layered alumino-silicate) that can form microscopic crystals called clay. Chemically it is hydrated sodium calcium aluminum magnesium silicate...
hydroxide \((\text{Na, Ca})_{0.33}(\text{Al, Mg})_2(\text{Si}_4\text{O}_{10})(\text{OH})_2 \cdot n\text{H}_2\text{O}\) and ions such as potassium and iron can exist as substituents. Montmorillonite forms layered structure. A number of cations and water molecules can exist in the interlayer region thereby making a mineral having high shrink-swell and cation exchange capacity.

The absorption peak at 3698 cm\(^{-1}\) was assigned to stretching frequency of structural O–H, of Kaolinite [47]. It was suggested that the peak helps to identify kaolinite from other clay minerals without the use of other complimentary techniques. We found this feature in all the samples and along with other frequencies (see Table 3) confirm the presence of kaolinite in the samples. Kaolinite is a clay mineral having low shrink-swell and cation exchange capacity. It is represented by molecular formula \(\text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4\). It is a layered silicate mineral in which tetrahedral sheets of silica (SiO\(_4\)) are linked through oxygen atoms to the octahedral sheet of alumina (AlO\(_2\)).

Illite is difficult to confirm in presence of other clay minerals. However, the presence of structural O–H stretching around 3620 cm\(^{-1}\) coupled with Al–Mg–OH deformation at around 830 cm\(^{-1}\) and Al–O–Si in plane vibration around 750 cm\(^{-1}\) may indicate illite [48]. These characteristic peaks along with other frequencies (given in Table 3) were present in all samples indicating illite. Illite is also a clay mineral that falls under the pylosilicate group of minerals \((\text{K, H}_3\text{O})(\text{Al, Mg, Fe})_2(\text{Si, Al})_4\text{O}_{10}[(\text{OH})_2(\text{H}_2\text{O})]\). It is non-expanding clay mineral having cation exchange capacity similar to kaolinite. Illite consists of sandwich structure of silica tetrahedron–alumina octahedron–silica tetrahedron layers.

Different carbonate minerals, such as polymorphs of CaCO\(_3\) (calcite, vaterite, aragonite), dolomite, thermonatrite are some of the commonly reported carbonate minerals in atmospheric dust samples [29, 30, 31, 32, 49]. Vaterite is one of the highly abundant polymorph of CaCO\(_3\). It can be identified in FTIR spectrum by the existence of out-of-plane bending mode.
near 876 cm\(^{-1}\) (\(\nu_2\)), in-plane bending mode near 745 cm\(^{-1}\) (\(\nu_4\)), stretching modes near 1087 cm\(^{-1}\) (\(\nu_1\)) and 1430 cm\(^{-1}\) (\(\nu_3\)), combined modes 1800 cm\(^{-1}\)(\(\nu_1 + \nu_3\)) and 2520 cm\(^{-1}\) (\(\nu_1 + \nu_2\)) [49–51]. The vibrational frequencies characteristic of vaterite were present in all samples.

Calcite is the next most abundant polymorph of CaCO\(_3\). It can be identified by the presence of in-plane bending mode near 715 cm\(^{-1}\) (other modes being same as of vaterite) [49, 50, 51]. The vibrational frequencies characteristic of vaterite were present in all samples. Dolomite can be identified by IR spectra by the existence of \(\nu_4\) mode at 730 cm\(^{-1}\) \(\nu_2\) mode at 882 cm\(^{-1}\), but the \(\nu_1, \nu_1 + \nu_4, \nu_1 + \nu_2\) modes overlap to that of calcite and vaterite [51]. These vibrational frequencies characteristic of dolomite were present in all samples. Dolomite is anhydrous mixed carbonate of calcium and magnesium and is represented by CaCO\(_3\), MgCO\(_3\).

Iron rich minerals are also reported in dust samples and hematite is one of the most abundant iron minerals having formula of Fe\(_2\)O\(_3\). This mineral can be identified by the Fe–O stretching frequency at 530 cm\(^{-1}\) and bending frequency at 452 cm\(^{-1}\) [52]. These frequencies were present in all samples indicating the presence of hematite in all samples.

Presence of asbestiform minerals in atmospheric dust samples was reported in several studies [29, 30, 33, 53]. Asbestos is the generic term used for naturally occurring crystalline silicate minerals having thin and long fiber with each fiber composed of many micofibrils. According to the physical make-up of the fiber, asbestiform minerals can be classified into serpentine and amphiboles groups. The serpentine group is characterized by curly fibers. Chrysotile (also known as white asbestos) falls in this group. The amphibole group of minerals contain pointed fibers and anthophyllite, crocidolite (also known as blue asbestos), actinolite, amosite (also known as brown asbestos), and tremolite fall in this group [54, 55]. It is well documented that inhalation of dust containing asbestiform minerals, depending on the expose dose and type, cause multitude of health issues, such as pleura, asbestosis, lung cancer, mesothelioma and other cancers [10, 56, 57, 58].

**Table 3. The vibrational frequencies observed in FTIR spectra of different samples (S1–S11) and assignment to the corresponding minerals.**

| Frequency (cm\(^{-1}\)) | Nature of band | Assignment |
|------------------------|----------------|------------|
| 3698                   | sharp peak     | stretching of structural O–H of kaolinite |
| 3688, 3691             | shoulder       | stretching of structural (surface) O–H of chrysotile |
| 3645                   | shoulder       | average envelope of stretching of structural (inner) O–H of chrysotile and O–H of crocidolite |
| 3620                   | sharp peak     | average envelope of O–H stretching of intercalated water of kaolinite, stretching of structural O–H of crocidolite, montmorillonite and illite, stretching of O–H of water bounded directly to cation (Fe\(^{3+}\) or Ca\(^{2+}\)) in the interlayer region of montmorillonite |
| 3550                   | broad peak     | O–H of surface bonded water in montmorillonite |
| 3405                   | broad peak     | O–H stretching of surface adsorbed water in clay minerals |
| 3280                   | shoulder       | bending overtone of surface adsorbed water |
| 2520                   | broad peak     | combination mode of calcite and vaterite |
| 2925                   | sharp peak     | C–H stretching (hydrocarbon) |
| 2850                   | sharp peak     | C–H stretching (hydrocarbon) |
| 1805                   | weak peak      | combination mode of calcite and vaterite |
| 1740                   | broad peak     | C–O stretching (organic compounds) |
| 1640                   | broad peak     | O–H bending (surface water) |
| 1450                   | broad peak     | stretching mode carbonate C–O (calcite, vaterite) |
| 1143                   | shoulder       | Si–O stretching (quartz) |
| 1087                   | shoulder       | average envelope of Si–O stretching of quartz and combination mode of calcite and vaterite |
| 1027\(^{*}\)           | broad peak     | average envelope of Si–O stretching of montmorillonite, kaolinite, illite, crocidolite, chrysotile |
| 914                    | weak shoulder  | deformation/bending vibration of structural OH of kaolinite, montmorillonite, illite, chrysotile |
| 876                    | weak peak      | out of plane bending of carbonate C–O of calcite and vaterite |
| 882                    | weak shoulder  | out of plane bending of carbonate C–O of dolomite |
| 777                    | broad peak     | silicate chain vibration of quartz and crocidolite |
| 750                    | weak peak      | Al–O–Si in plane vibration of illite, |
| 745                    | weak peak      | in plane bending of carbonate C–O of vaterite |
| 730                    | weak shoulder  | Al–Mg–OH deformation of illite, in plane bending of carbonate C–O of dolomite |
| 715                    | weak peak      | in plane bending of carbonate C–O of calcite |
| 694                    | weak peak      | silicate ring vibration of quartz, kaolinite, and crocidolite |
| 670                    | weak peak      | silicate ring vibration crocidolite |
| 652                    | broad weak peak| average envelope of silicate ring vibration of crocidolite and vibration of outer Mg–OH of chrysotile |
| 605                    | weak shoulder  | vibration of inner Mg–OH of chrysotile |
| 541, 505               | weak shoulder  | cation oxygen stretching of crocidolite |
| 530                    | broad peak     | Fe–O stretching of hematite |
| 524                    | weak shoulder  | Al–O–Si deformation montmorillonite and illite |
| 471                    | weak shoulder  | Si–O–Si deformation of illite and chrysotile, bending vibration of Mg–O bond of chrysotile |
| 452                    | weak peak      | Fe–O bending of hematite |
| 460                    | broad peak     | average envelope of Si–O–Si deformation of quartz and montmorillonite |
| 436                    | broad peak     | Si–O–Si deformation of illite and chrysotile, bending vibration of Mg–O bond of chrysotile |
| 425                    | weak shoulder  | Si–O–Si deformation of kaolinite |

\(^{*}\)spectral de-convolution shows this peak around 1010 cm\(^{-1}\).
addition, outer Mg–OH vibration frequency at 650 cm$^{-1}$, inner Mg–OH vibration at 604-611 cm$^{-1}$, are the other important frequencies that helps to identify chrysotile form other minerals [59, 60, 61]. These characteristics frequencies in FTIR measurements were present in all dust samples thereby indicating the presence of the asbestiform mineral chrysotile. Chrysotile is a fibrous, soft and disordered asbestos of serpentine group and commonly represented by a chemical formula of Mg$_6$(Si$_2$O$_5$)(OH)$_$_$_4$. It is one of the most commonly used asbestos and its prolonged exposure is reported to cause lung cancer.

Crocidolite, on the other hand, can be identified by the presence of stretching of the structural O–H at 3645 and 3620 cm$^{-1}$, Si–O stretching vibration at in the range of 990–1143 cm$^{-1}$, silica chain and ring vibration at 778, 725, 694 cm$^{-1}$, and cation-oxygen stretching vibration at 541 and 505 cm$^{-1}$ [59]. These characteristic frequencies were present in samples S6–S11. Crocidolite is reported to be the most toxic form of asbestos. It is the fibrous form of the amphibole mineral riebeckite with chemical formula of Na$_2$(Fe$^2$)$^3$(Fe$^3$)$^2$Si$_8$O$_22$(OH)$_$_$_$_2$.

The organic carbon can be detected from the presence of C–H stretching frequency at 2925 and 2850 cm$^{-1}$. The organic carbon was detected all the samples. This is expected observation as the sampling was done in the densely populated and vehicular area of Kathmandu.

A summary of the vibrational frequencies observed in FTIR spectra of different samples and their assignment to the corresponding minerals is shown in Table 3.

### Table 3.

| Minerals            | Samples | 2θ (°)   | dhkl (Å)  |
|---------------------|---------|----------|-----------|
| Illite              | S1–S11  | 17.8, 23.7, 22.08, 26.3, 28.90, 31.3, 45.90, 50.36 | 4.99, 3.76, 3.86, 3.39, 3.09, 2.86, 1.98, 1.81 |
| Quartz              | S1–S11  | 17.8, 20.87, 24.3, 26.70, 36.06, 36.60, 39.50, 40.36, 42.51, 45.90, 50.20, 54.94, 55.38, 55.6, 60.0, 67.79, 68.00, 68.18, 68.37, 73.51, 75.70, 77.70 | 4.93, 4.26, 3.68, 3.34, 2.49, 2.45, 2.28, 2.23, 2.13, 1.98, 1.82, 1.67, 1.66, 1.65, 1.54, 1.38, 1.38, 1.377, 1.274, 1.29, 1.25, 1.23 |
| Kaolinite           | S1–S11  | 12.5, 17.80, 19.86, 23.13, 40.36, 45.90, 50.66, 55.7, 61.6, 64.06 | 7.15, 4.99, 4.49, 3.85, 2.23, 1.98, 1.80, 1.65, 1.50, 1.45 |
| Montmorillonite     | S1, S2, S3, S4, S5, S6, and S11 | 19.8, 27.2, 35.03 | 4.49, 3.28, 2.56, 1.50 |
| Crocidolite         | not detected |                        | 7.30, 4.50, 3.65, 2.44, 1.53 |
| Chrysotile          | S1–S11  | 12.1, 19.7, 24.3, 36.7, 60.1 | 3.85, 3.07, 2.49, 2.28, 2.09, 1.91, 1.87 |
| Vaterite            | S1–S11  | 20.7, 25.2, 27.5, 32.2, 43.4 | 4.27, 3.54, 3.24, 2.78, 2.08 |
| Dolomite            | S1–S11  | 24.2, 31.42, 49.5, 50.6, 51.2 | 3.68, 2.88, 2.19, 1.84, 1.80, 1.78 |
| Hematite            | S1–S11  | 19.9, 22, 24.3, 33.3, 35.7, 37.5 | 4.46, 4.04, 3.68, 2.69, 2.50, 2.34 |

Majority of intense peaks in XRD data represented quartz mineral, which is due to high crystallinity order of quartz. One of the approaches to measure the crystallinity of quartz is to measure the crystallinity index (CI) in the scale of 0–10; 0 referring to the amorphous material and 10 referring to the perfectly crystalline material [38]. The presence of five diffraction peaks in the range of 20 value of 67° to 69° indicated crystallized quartz [38]. The XRD pattern in the region of interest for samples S5 and S11 is shown in Figure 8A. We found quintuplet peaks indicating the presence of quartz of high crystalline order. The CI values of the dust samples ranged from 7 to 9.7 (Figure 8B). This indicated that in all samples studied silicate mineral of high degree of crystallinity was present.

#### 4.4. Comparison of different data

The SEM-EDX data are consistent with XRD and FTIR data. For example, sample S4 consisted of minerals such as calcite, dolomite, vaterite, ilite, montmorillonite, kaolinite, chrysotile, hematite, and quartz. The types of elements required for the formula unit of these minerals are also found to exist in the EDX data (refer to Table 2 for sample S4). It is interesting to note that the single particle data for samples S2 and S3 (S2-1, S3-1, S3-2) contained different elemental composition than EDX spectrum collected for whole field of view (S2 and S3 in Table 2). The single particle spectrum S2-1 contained Ca, Si and Al as the metallic element indicating that the particle could be rich in calcite, vaterite, and quartz.

#### 4.5. Source of minerals in dust

In Kathmandu Valley, almost all structures are concretized. In this respect, cement can be one the sources of the minerals in the particulate matter. To explore this, we measured XRD data of three cement samples obtained from three major cement suppliers. The samples looked very similar in XRD and a representative datum along with sample 11 (S11) is plotted in Figure 9. For clarity, the data are plotted in the range of 20 value of 10°–26° in frame A, 26°–53° in B, and 53°–79° in frame C. Interestingly, the cement samples also showed peaks characteristics of: i) crystalline silica (quartz), ii) carbonate minerals vaterite, calcite, and dolomite, iii) asbestiform mineral chrysotile. The cement samples also showed dominant peaks characteristics of calcium hydroxide, calcium silicate, and iron rich mineral ferrite (rich in hematite). These minerals are also reported in cement samples of other countries [70]. This observation indicates that cement could be one of the sources of quartz, vaterite, calcite, dolomite, and chrysotile minerals in the dust sample studied.

The earlier studies reported that the cardiovascular related diseases and deaths are increasing in Nepal [24, 71]. It is established fact that air...
pollution is one of the major contributors to respiratory and cardiovascular diseases [6, 72]. Our study showed that dust samples collected from the densely populated locations of Kathmandu Valley contain different minerals including crystalline silica (quartz) and asbestiform minerals chrysotile and crocidolite. Inhalation of crystalline silica and asbestos rich air cause different pulmonary and cardiovascular diseases and eventually to death. Without the information on concentration of minerals, exposure dose, and epidemiological studies, it is not possible to comment on the cardiovascular related health issues, however, we believe that the findings of this work will be important to understand the respiratory and cardiovascular related health issues in Kathmandu Valley.

Furthermore, we also like to mention the morphological and chemical analysis of particulate matter samples from different Asian cities other than Kathmandu. Study on the air samples collected from different cities of Saudi Arabia reported that particles having rod, irregular, and circular shape existed in the dust samples [73]. A study made in air dust samples of Agra, India showed the presence of mineral, soot, fly ash, tarballs, aluminosilicates/silica, fluorine, carbon rich, and Cl-Na rich particles [74]. A similar study in samples collected from rural and urban area of Delhi [75] and Lucknow [76] was also reported. Morphology and composition of particulate matter collected from Peshawar, Pakistan was made by Zeb et al [77]. In the study particles, based on morphology and composition, were classified as: aluminosilicates (23%), silica (12%), carbonaceous (49%), calcium rich (3%), chloride (2%), Fe/Ti oxides (3%), sulfate (5%), biogenic (3%). A long term study (2012–2014) on PM2.5 samples collected from twenty urban cities of China was reported by Liu et al [78]. It was found that the major PM2.5 constituents across all the urban sites were organic matter (26.0 %), elemental carbon (6.0 %), SO4−/C0 (17.7 %), NO3−/C0 (9.8 %), NH4+ (6.6 %), Cl− (1.2 %), mineral dust (11.8 %), and unaccounted matter (20.7 %).

5. Conclusions

To summarize, we identified different minerals in dust samples collected from eleven core urban areas of Kathmandu Valley. We found carbonate minerals (calcite, vaterite, and dolomite) iron rich mineral (hematite), clay minerals (illite and kaolinite), crystalline silicate mineral (quartz), and asbestiform mineral present in all samples. The clay minerals montmorillonite and asbestiform mineral crocidolite were found in seven and five samples, respectively. This observation was supported by EDX and XRD measurements. Electron microscopic measurements showed particles having very diverse morphology and very few particles of asbestiform type. The crystallinity index (CI) of quartz mineral in the samples ranged from 7 to 9.7 indicating that quartz of high crystallinity present in all the samples. The major elements in the dust samples were found to be C, O, Mg, Ca, and Si. The XRD data analysis in three different brands of cement samples showed the presence of crystalline silica (quartz), carbonate minerals (vaterite, calcite, and dolomite, and

Figure 8. (A) XRD data for S5 and S11 plotted in the range of 20 values of 67° to 69°. (B) The calculated crystallinity index of all the samples.

Figure 9. XRD pattern of cement sample (gray color) and sample 11 (black color). For clarity, the data are plotted in the range of 20 value of 10°–26° in frame A, 26°–53° in B, and 53°–79° in frame C. For easy comparison the S11 peaks ~21° and 26.7° are truncated. Ch = chrysotile, Q = quartz, C = calcite, CS = calcium silicate, CaH = calcium hydroxide, D = dolomite, F = ferrite, and V = vaterite. The peaks that closely overlap in two samples are labeled in black color and the intense peaks present only in cement samples are labeled gray color.
asbestiform mineral chrysotile; indicating that cement could be one of the sources of minerals in dust samples.

Declarations

Author contribution statement

Bhanu Neupane: Conceived and designed the experiments; Analyzed and interpreted the data; Wrote the paper.

Amita Sharma, Mahesh K. Joshi: Performed the experiments; Analyzed and interpreted the data.

Basant Giri: Analyzed and interpreted the data.

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The authors declare no conflict of interest.

Additional information

No additional information is available for this paper.

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