Sampling of Atmospheric Precipitation and Deposits for Analysis of Atmospheric Pollution

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This paper reviews techniques and equipment for collecting precipitation samples from the atmosphere (fog and cloud water) and from atmospheric deposits (dew, hoarfrost, and rime) that are suitable for the evaluation of atmospheric pollution. It discusses the storage and preparation of samples for analysis and also presents bibliographic information on the concentration ranges of inorganic and organic compounds in the precipitation and atmospheric deposit samples.

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1. INTRODUCTION

The tremendous dynamics of the atmosphere makes it the main propagation path for air pollution and its transport between the various elements of the environment in the form of dust, gases, and aerosols. Pollution, depending on its properties and on meteorological conditions, is subject to scattering and transformation during atmospheric transport. Most of the pollutants eventually return to the earth’s surface, often at great distances from their sources. Their return may be through precipitation or absorption of gaseous pollutants and aerosols by surface waters, the vegetation cover, or the soil. Of these, wet deposition plays the greatest role in feeding atmospheric pollutants back to the earth’s surface in regions distant from their sources.

Atmospheric precipitation is the result of condensation of atmospheric water vapour due to adiabatic cooling. Precipitation falls down to the surface in the form of rain, snow, drizzle, snow pellets, and hail; floats in the air in the form of clouds and fog, and settles on surfaces in the form of dew, hoarfrost, and rime (see Figure 1).

Environmental protection begins with identification and definition of the kind and degree of pollution and its analysis and monitoring. During the past 20 years the field of analysis and monitoring of atmospheric deposition has grown dramatically because we now recognize that the quantity and quality of transported pollutants, as well as the range of their interactions, can be a good indicator of the degree of atmospheric pollution in a given geographical region. An extensive review of the design and basic parameters of samplers for rain precipitation and runoff waters appeared in 2002 [1]. This paper is prompted by the growing interest in the sampling of other atmospheric deposits, particularly fog and clouds, dew, hoarfrost and rime, that began around 1990. The basic characteristics of the forms of precipitation that will be discussed in this paper are presented in Table 1.

2. IMPORTANCE OF REPRESENTATIVE SAMPLES

To be a source of reliable analytical information a sample must be representative of the object or distribution from which it is extracted. In practice the samples we analyze can only be a small fragment of the objects of interest. Hence, we must take care to insure that the samples are as representative as possible [2]. Sampling has a special significance for all analytical processes; in a sense, it is the critical point of analysis. Errors committed at this stage cannot be estimated nor can their effect upon the result of the analysis be reduced.

One of the most important ways in which sampling can fail to be representative is through sample contamination. For sampling of atmospheric deposits, the proper preparation of the sampler is an important factor for minimizing sample contamination. The most obvious step is also the most important—cleaning the sampler before collecting a sample. A standard cleaning procedure includes, among other things, washing in deionized water [3–5] and/or for example, rinsing with acetone [6]. Collecting vessels are also washed in water with detergents [7], in aqueous solutions of nitric acid [7], distilled water, and eventually rinsed with deionized water [8, 9]. Subsequent handling of samples in...
and automate, and should require minimal maintenance. With any instrument, the collector should also be easy to use for large amounts of liquid water for wet chemical analysis. As stages of collection, and provide rapid collection of relatively efficient, quantitative water collection, but must also avoid collection of submicron “nonactivated” aerosols, preserve the size and chemical composition of droplets through all the stages of collection, and provide rapid collection of relatively large amounts of liquid water for wet chemical analysis. As with any instrument, the collector should also be easy to use and automate, and should require minimal maintenance.

3. SAMPLERS FOR THE COLLECTION OF FOG AND CLOUD WATER SAMPLES

As previously noted, transport of anthropogenic pollutants through the atmosphere is an important means of their worldwide distribution. Airborne pollutants can be transferred to aquatic and land environments via mechanisms such as wet and dry deposition and air/water partitioning. Fog has recently attracted the attention of the scientific community as a potentially important deposition mechanism. Organic and inorganic pollutants distribute themselves between the vapour and aqueous phases, of fog and between the particles present in both phases and chemical reactions can occur within fog droplets. When fog droplets subsequently evaporate, the chemicals contained in them can be deposited on surfaces in contact with the fog [10].

Over the last 20 years, investigators have examined the chemical compositions of cloud and fog [7, 11-13]. They have studied the processes occurring in atmospheric particles and developed sample collection methods [14, 15]. Collecting samples of fog and cloud water is often more complicated than collecting samples of precipitation or runoff water [1], since we need to extract the water from the cloud or fog. Many of the methods for fog and cloud water sampling are based on instruments originally designed for quantitative meteorological measurements of the amount of water in clouds or fog. Fog and cloud water samplers that are used for the evaluation of pollutants differ from those used for water measurements, because they must not only perform efficient, quantitative water collection, but must also avoid collection of submicron “nonactivated” aerosols, preserve the size and chemical composition of droplets through all the stages of collection, and provide rapid collection of relatively large amounts of liquid water for wet chemical analysis. As with any instrument, the collector should also be easy to use and automate, and should require minimal maintenance.

Cloud and fog water collectors operate primarily on the principle of inertial impaction on a plane surface, a standard technique also used for the collection of dry aerosol particles. Collection efficiency curves, which show the percentage of particles of any size which are collected as a function of the particle size, indicate that the instruments based on inertial impaction generally provide more reliable results than other methods [16].

The efficiency curves of inertial impactors show a sharp division (cutoff) between the droplets collected and those which are not. Marple and Willeke [17] formulated design criteria for inertial impactors that enable construction of dry aerosol impactors with well-defined cutoff characteristics. Their techniques had to be adapted, however, to the special needs of liquid water sampling [16]. Single stage cloud water impactors, based on the adapted techniques, have been used for several years to perform studies of cloud and fog water chemical composition [16]. They are used in both active collectors, where flow of air containing the droplets is forced by means of a suitable mechanical device, and passive collectors, where natural circulation of air (wind) is utilized. There is a wide variety of passive and active collectors available for various ambient conditions.

4. PASSIVE COLLECTORS

Passive collectors are often simpler in operation and may be used in windy environments. However, their impaction characteristics (cutoff) are less controllable. The collecting elements (impaction plane) for passive collectors may vary from flat surface to solid elements like rods; tubes as well as strings, ropes, filaments, screens, and meshes have also been used.

The simplest passive fog sampler is the deposition plate, typically a horizontal plate on which fog droplets are allowed to settle [18]. This sampler may suffer from contamination due to dry deposition and dew formation, which leads to significant biases toward errors with large particle measurements. The design is attractive on its own, but its disadvantages restrict its applications.

The next simplest fog sampler is the string screen sampler in which fog is collected through the impaction of fog droplets on a string screen. After collision the droplets adhere to the string and drop along the strings to a collection tray. String screen samples can be also in an active version.

In a published paper [19], a passive collector is described (Figure 2), consisting of a 2 m tall collection, of two horizontal disks 20 cm in diameter, installed vertically on frames at a distance of 40 cm from each other. Between the disks, Nylon strings of 0.2 mm diameter are stretched into two rows. The fog water collected on the strings is stored in a 500 mL polyethylene bottle. Collection area was 314 cm². A hood above the collector prevents rain from diluting the fog sample. The sampler was set up in a 1 m high PVC tube to protect it from direct radiation and light.

The principle of the work of the collector described in [20] (Appalachian Mountain Club/Worcester Polytechnic Institute (AMC/WPI)) is based on the utilisation of wind to transfer particles to Teflon collecting strands. The cloud
Table 1: The characteristics of some atmospheric precipitation types.

| Atmospheric precipitation or deposits | Characteristics |
|--------------------------------------|----------------|
| Cloud                                | Water droplets of microscopic size (diameters not exceeding 100 µm) and/or ice crystals floating in the air due to microturbulences counteracting their gravitational sedimentation. |
| Fog                                  | Suspension of very small water droplets with diameters below 0.05 mm or ice crystals in air. It is formed from the result of cooling from the ground layer of air to the dew–point temperature–reaching the state of saturation with water vapour. From the sample collection point of view both cloud and fog present similar challenges. |
| Dew                                  | Deposit of water droplets forming on the surface of ground and objects on earth’s surface or close to it due to condensation of water vapour contained in surrounding air. Generally, it is caused by night-time radiation of heat. Dew originates also when warm and humid airflows above a cooled-down substratum, whose temperature is lower than the saturation point of the inflowing air mass. |
| Hoarfrost                            | Deposit of ice crystals forming generally on horizontal surfaces. The appearance and size of the deposit depend on thermal properties of the substratum and on air humidity. It occurs with distinct differences between air and ground temperatures and the ground surface temperature must be negative but the air temperature can be positive. It is often linked with ground frost. |
| Rime                                 | Granular or crystalline deposit (generally both forms of deposit appear simultaneously) settling most easily on thin fibrous objects windward, at negative air temperature and with fog. The amount of the deposit depends on the density and duration of fog. |

Figure 2: Structure of a passive fog water collector [19].

Figure 3: Structure of a passive cloud water collector (AMC/WPI) [20]: (a) top view, (b) side view.

Water droplets are collected principally by the mechanism of inertial impaction on Teflon strings. Exclusion of the heavier rain droplets is accomplished by both the placement of the collection strands deep within the collection box and by a baffle system (Figure 3). The airflowing through the baffles is restricted and forced to turn, causing it to accelerate. The rain droplets, with their greater inertia, overcome the viscous drag effects of the airstream and pass out of it, impacting on the lower baffle. A drain below the baffle permits the separated rain water to leave the collector. The cartridge containing the collection strands is located behind the baffle to allow adequate expansion of the airflow and thus to maximize the utilization of the collection surface area. The distance back from the top of the entrance of the collection box to the upper lip of the lower baffle was selected for its theoretical ability to remove free-falling droplets > 200 µm at winds of 0–10 m/s and droplets > 500 µm at winds of 0–25 m/s. These values represent the minimum ability of the collector to exclude rain and drizzle. To prevent
contamination of the cloud water sample by rain, baffles have lips to prevent the impacted rain water from running to the edge of the baffle and from becoming reentrained in the accelerated airflow. The upper baffle has a small reservoir where water collects prior to draining out holes drilled on the side of the collector. To prevent rain splatter on the lower surface of the collector from reentering the airstream, a series of vanes set at 40 degrees is positioned in front of the lower baffle. Rain water collected by these vanes leaves the collector at the base of the lower baffle.

In order to collect cloud water samples, another collector has also been used, consisting of 0.45 mm diameter Teflon-coated wires, strung at 3 mm intervals around the perimeters of two 25 cm diameter plastic disks, held 1 m apart by plastic rods [21]. The surface area of the collector is sufficient to provide a sample of 50 mL in 3–30 minutes, depending on the wind speed and cloud liquid water content. The collection efficiency of this sampler indicates that cloud droplets of 10 µm diameter are collected with only 50% efficiency at wind speed of 1 m/s, proving poor performance of the collector at low wind speeds. For 80% of the cloud water samples, the wind speed was greater than 5 m/s, yielding 50% collection diameters below 5 µm.

5. ACTIVE COLLECTORS

Active collectors use either forced flow (fans, pumps) or motors moving the collecting elements in the air (usually rotating them) to achieve the same end. Active collectors typically use collection elements such as rods, tubes, strings, ropes, filaments, screens, and flat surface. Some active collectors use jet-driven impaction onto solid surfaces. Size-resolved cloud composition is usually obtained via active collectors with multiple jet/impaction surface combinations, or multiple stages with varying cutoff diameters.

The active string screen fog sampler described by Jacob et al. [22] consists of three parts: a series of three screens of Teflon wires on which the fog condenses, a baffle which smoothes the airflow, and a fan which pulls the air past the Teflon wires. The fog water is collected on the wires until drops are formed, then they move down the wire, pool in a Teflon tray, and are then collected in a clean glass jar [10]. Inclining the screen at 35 degrees from the vertical in the direction of airflow helps prevent resuspension of impacted droplets into the airflow. During sampling the fog water only comes into contact with Teflon and Glass. During sample collection, the face velocity of the air through the sampler, the time of collection, and the air temperature are monitored. Typical sample collection volumes range from 50 to 200 mL, which can take from 50–180 minutes depending on the liquid water content LWC of the fog event (0.1–0.3 g/m³) [23]. Droplets in the range 3–100 µm diameter are efficiently collected using this type of sampler [22].

Figure 4 presents a schematic diagram of a typical active string screen collector. The Teflon screen consists of a frame of four 4 mm thick copper rods, over which Teflon strings are strung [24]. Drops collected on the strings find their way to polyethylene collecting bottles. Additional string screen collectors are described by Sasakawa and Uematsu [25] and Sasakawa et al. [26].

The High-Volume Fog Sampler [27, 28] is a scaled-up version of the sampler described by Jacob et al. [22]. In this system, a 50 cm diameter fan in the back draws air at the rate of 4400 m³/h across a screen consisting of four layers of 0.28 mm Teflon filaments wound around threaded rods. Fog droplets impact on the Teflon filaments, coalesce, and flow down the filaments into a Teflon-coated funnel. The fog water then drains by gravity through a Teflon tube to a Teflon bottle. Collection rate of this system is approximately 1 L/h in fog with 400 m visibility.

The Active String Cloud Water Collector (CWP) described by Daube et al. [20] is shown schematically in Figure 5. This system collects cloud droplets on a removable cartridge of 0.78 mm diameter Teflon strands. A fan inside the collector draws air and cloud droplets up through a ventral opening and then into the vertical collection strands. The positioning of the air inlet on the bottom of the sampler makes it possible to avoid collecting rain [29–31]. The collector excludes most rain droplets ≥ 200 µm at wind speeds ≤ 10 m/s. With this system, cloud water collection is usually initiated within 15–30 minutes after the onset of a cloud event, and the typical collection time is approximately 5 hours.

The CalTech Active Strand Cloud Water Collector (CASCC), built at California Institute of Technology, is shown schematically in Figure 6. It has been described in detail in several publications [4, 11, 32–35]. In this system, cloud droplets are collected by inertial impaction on an angled bank of six rows of 508 µm diameter Teflon strands. A fan sucks in air through the Teflon strings with a velocity of 8.5 m/s. The strands are inclined at an angle of 35 degrees from vertical. The collected droplets coalesce, and are drawn down the strands by gravity and aerodynamic drag into a Teflon trough. A Teflon tube delivers the sample from the trough to a collection bottle, which is emptied at 30–60 minutes intervals. The 50% collection efficiency cutoff, based on droplet diameter and predicted from impaction theory, is 3.5 µm. A protective rain shield, with its opening facing downward, can be attached to the front of the collector to
exclude large sedimenting droplets \( (d > 300 \mu m) \) [34]. The typical flow rate of air through the CASCC is 24.5 m³/min, yielding a collection rate of approximately 2 mL/min when the LWC of the fog is 0.1 g/m³.

The CASCC cannot function below 0°C, because cloud water droplets will freeze on the collection surface. For this reason, a winter cloud water sampler—the CalTech Heated Rod Cloud Water Collector (CHRCC)—was developed [36]. The stainless steel rods that form the collection surface in this sampler are internally heated on a periodic basis when temperatures fall below 4.5°C (Figure 7). When heated, accumulated frozen cloud water on the rods melts and drains off the rods to the sample bottle. A fan draws air across an inclined bank of six rows of the 3175 μm diameter rods at a rate of 6.3 m³/min. The corresponding cutoff of the rod bank is calculated to be 7.7 μm. The predicted collection rate in a cloud with an LWC of 0.1 g/m³ is 0.44 g/min.

6. MULTISTAGE FRACTIONING SAMPLERS

The operation of Two-Stage Fog Water Impactor (TFI) [16] is also based on the principle of inertial impaction on a plane surface. This sampler consists of vertical slit impaction stages, one to collect the larger droplets, followed by two identical stages in parallel. These collect those droplets which passed the first stage, but which are above a well-defined cutoff diameter. The cutoff diameters of the first stage are between 10 and 12 μm, and of the second stages are between 5 and 6 μm, calculated for flow rates from 150 to 200 m³/h. The air, together with smaller water droplets, leaves the impactor through holes in the plate and four suction pipes. The collected samples are forced by airflow to the external edges and to streams directed down the plates. The samples are collected into vials situated at the bottom of the device. This apparatus has the capability of controlling the velocity of incoming air depending on the average wind speed.

Cloud chemistry can vary as a function of drop size. In order to investigate variations in chemical composition across the drop size spectrum, a multistage cloud water collector was developed by Moore et al. [37] at Colorado State. The CSU 5-Stage Collector (Figure 8) is a cascade inertial impactor that collects samples of cloud water in five independent size fractions for chemical analysis. Its design incorporates many features to facilitate its use in the field, and to maintain both consistent performance between varying atmospheric conditions and the chemical and physical integrity of the collected sample. The sampler consists of five stages, each with a single, one-sided rectangular jet arranged
in a cascade. The intensity of airflow rises as it passes through the sampler. Drops of progressively smaller diameters are collected in each stage as those with too much inertias cannot follow the fluid streamlines and impact. The collector is mounted at 45 degrees to the horizontal, so sampled drops coalesce and run down to polypropylene vials threaded directly into each stage. The collector is oriented into the wind during operation, subject only to site restrictions and its own geometry. While the collector is designed for low wind environments, a baffle or windshield parallel to the inlet can be added for higher winds. The experimentally determined virusments, ab aba geometry. While the collector is mounted at 45 degrees to the horizontal, so sampled drops do not follow the fluid streamlines and impact. The collector is mounted into the wind during operation, subject only to site restrictions and its own geometry. While the collector is designed for low wind environments, a baffle or windshield parallel to the inlet can be added for higher winds. The experimentally determined virusments, ab aba geometry.

Warm, dry, and clean air

Suction

Stagnation plane

Droplets of different size (inertia)

Inner tube

Porous section

Figure 9: A schematic diagram of CVI sampler.

Dry air inside the CVI sampler. The maximum droplet radius that can be sampled is between 50 and 100 µm. Ambient gases and submicron aerosol particles are rejected in the CVI with almost 100% effectiveness. The instrument requires a condenser for water collection and measurement. In this respect, a CVI collecting sample for chemical analysis must ensure total recovery from the gaseous phase of both water and other volatile substances (pollutants) contaminating original droplets caught by the device.

7. ROTATING COLLECTORS

The active sampler described by Glotfelty et al. [6] is a rotating screen device, 50 cm in diameter, in which four layers of stainless steel screen are rotated around a central axis at 720 rpm. Fog water obtained from droplets impacting on the screen is centrifuged to the periphery, collected in a slotted aluminum tube, and drained into a collection vessel. A large fan pulls air through the device at 160 m³/min. Under these conditions, the collection rate of this system is approximately 0.5–1 L/h depending on the LWC of fog water (0.024–0.08 g/m³).

The CalTech, Atmospheric Science Research Center (ASRC), and AeroVironment (AV) instruments are rotating collectors [14], employing external surfaces for the impaction of the droplets. Jacob et al. [39] built the CalTech system. It is an external impactor that sweeps through the air at a high velocity (1700 rpm) in order to collect large particles. The arm (63 cm long) spins in a vertical plane, driven by a motor (Figure 10). Each end of the arm has a slot milled into its leading edge. Bottles (30 mL) are mounted at the ends of the arm to collect the water that impacts on the slots. Threaded Teflon tubes are screwed on the ends of the arm and extend inside the collection bottles, preventing the collected fogwater samples from running out after the instrument has stopped. Deflectors prevent water that impacts on the solid part of the arm from entering the slot. Small fins are welded to the back of the arm for extra strength. The entire arm is Teflon-coated to prevent chemical contamination and to facilitate cleaning.

Figure 10: Structure of a fog and cloud water rotating arm collector [11].
The rotating arm collector samples air at a rate of 5 m$^3$/min. Laboratory calibration indicates a lower size cutoff of 20 µm diameter (50% collection efficiency). This sampler has been applied in several studies [40, 41].

The ASRC sampler consists of 1 500.41 mm diameter Nylon strings mounted between two plates (Figure 11). The sampler rotates to its vertical axis at 100 rpm [14]. Water impacting on the strings collects in traps on the bottom plate. Periodically, the sample rotation is stopped and the fogwater on the strings is coaxed into the traps by tapping the bottom plate with a mallet. At the end of the sampling period, water in the trap is manually transferred to polyethylene bottles. Two versions of the device exist; one of them is passive and the other is active.

The AV rotating rod sampler collects droplets by impaction on a Teflon-coated rod rotated in a vertical plane at 3450 rpm (Figure 12). The outer part of the rod is 1.6 mm and the inner part is 19 mm in diameter to provide size cuts of 2.5 and 10 µm, respectively, [14]. Water impacting on the rods is transferred by centrifugal force to circular polyethylene troughs that drain to polyethylene collection bottles. Separate troughs and sample bottles are used for the two size fractions.

In Table 2 parameters of samplers used for collecting fog and cloud were set together.

### 8. OTHER TYPES OF SAMPLERS

The Global Geochemistry Mesh Impaction Fog Sampler and the Desert Research Institute (DRI) instruments are internal collectors [14], in which air is drawn into the instrument and extracted by surfaces internal to the device.

The Global Geochemistry Mesh Impaction Fog Sampler (Figure 13) is an internal impaction sampler that collects fog water on a 10 cm diameter by 4 cm thick polypropylene mesh located at the entrance of a V-shaped Teflon lined PVC pipe. The mesh is made of interlaced filaments (410 µm diameter) and has a void volume of 96%. Air is drawn through the mesh at 1.7 m$^3$/min. Fog droplets impact on the mesh, coalesce, and then drain into a polyethylene bottle at the bottom of the V-tube. The sampler can effectively intercept droplets > 5.0 µm, with a 50% collection efficiency at 2.4 µm. Liquid holdup on the mesh depends on the mass of liquid sampled. If ≤ 1 g of water is sampled, all of it remains on the mesh. If 100 g is sampled, less than 5% remains [14].

The DRI (Figure 14) is based on a jet impaction principle. Fog is drawn through three rectangular jets at a total flow rate of 20 L/s. The accelerated droplets impact on rotating Teflon rollers and are transferred to a central roller. Here, the fogwater is forced to accumulate in bulk form and is deposited into a polystyrene collection vessel. The impactor has a sharp cutoff at 5 µm diameter to allow efficient collection of droplets while rejecting small interstitial particles. The collector is housed in a shelter consisting of an inverted, insulated 2501 drum to prevent collection of precipitation. Airflow up to the collector is provided by a fan [14, 44].

Specially [8] designed equipment, consisting of a stainless steel cooling chamber (10 cm × 10 cm × 22 cm) and a collector for fog droplets utilizing the impaction technique, has also been used for collecting fog water. Fog air is drawn at the rate of 0.2 m$^3$/min and the fog droplets impact on the collector which is maintained at 15°C in the cooling chamber. The sampling period varies from 30 to 120 minutes, and in this time from 5 to 30 mL of fog water is collected. The volume of the sample collected depends on the duration of the fog event.

In the available literature [15, 46] one can find information on an electrostatic precipitation method capable of sampling single cloud or fog drops (Figure 15). The precipitator is based on the corona discharge principle. A copper electrode is placed as a discharge electrode at a distance of 10 cm above an aluminum precipitation electrode. A 25–30 kV voltage is applied to the discharge electrode for 1 second to produce a spray of electrons or negative charge and the charge is transferred to the droplets by the action of the electric field. Charged droplets moving in the direction of the collecting electrode are captured on Petri dishes. The droplets are covered immediately with paraffin oil to prevent evaporation and contamination. Capillary electrophoresis is then used for the chemical analysis of the individual drops. This sampler can also be used for collecting bulk phase cloud or fog water, by increasing the time for the application of the voltage to the discharge electrode from 1 second to 5–15 minutes, depending on the density of the fog.
| Name of sampler | Type of sampler | Collector surface | Collection rate (mL/min) | Cutoff (µm) | LWC (g/m³) | Flow rate (m³/min) | Comments | Reference |
|-----------------|-----------------|-------------------|--------------------------|------------|-----------|------------------|----------|----------|
| String screen sampler | Active | Teflon wires | 50–200/ 50–180 | 3–100 | 0.1–0.3 | | Inclining the screen at 35 degrees from the vertical in the direction of airflow helps prevent re-suspension of impacted droplets into the airflow. | Jacob et al. [22] |
| High-Volume Fog Sampler | Active | Teflon filaments | 16.7 | | | 4400 | 1 L/h in fog with 400 m visibility, 50 cm diameter fan. | Schomburg et al. [27], Chernyak et al. [28] |
| CWP | Active | Teflon strands, 0.78 mm diameter | | | | | The collector excludes most rain droplets ≥ 200 µm at wind speeds ≤ 10 m/s. | Daube et al. [20] |
| CASCC | Active | Teflon strands, 508 µm diameter | 2 | 3.5 | 0.1 | 24.5 | The strands are inclined at an angle of 35 degrees from vertical. | Igawa et al. [4, 33], Munger et al. [32, 34] |
| CHRCC | Active | Stainless steel rods | 0.44 | 7.7 | 0.1 | 6.3 | The stainless steel rods that form the collection surface in this sampler are internally heated on a periodic basis when temperatures fall below 4.5°C. | Collett et al. [36], Schell et al. [16] |
| TFI | Active | | 5–12 | | | 150–200 | This apparatus has the capability of controlling the velocity of incoming air depending on the average wind speed. | Schell et al. [16] |
| CSU 5-Stage Collector | Active | | 4.5–29 | | | | The collector is mounted at 45 degrees to the horizontal. It is a cascade inertial impactor that collects samples in five independent size fractions. | Moore et al. [38] |
| CalTech Rotating Arm Collector | Rotating active | Teflon-coated | | 20 | | 5 | The arm spins in a vertical plane, driven by a motor at 1700 rpm. | Jacob et al. [39, 41], Johnson et al. [40] |
| Rotating screen sampler | Rotating active | Stainless steel screen | 8.3–16.7 | 0.024–0.08 | | 160 | Fifty-centimeter diameter screen rotated around a central axis at 720 rpm. | Glotfelty et al. [6] |
| AV | Rotating active | Teflon-coated rod | | | | | Rod rotated in a vertical plane at 3450 rpm. | Hering et al. [14] |
| ASRC | Rotating | Nylon strings, 0.41 mm diameter | | | | | The sampler rotates to its vertical axis at 100 rpm. Two versions of the device exist, one of them is passive, and the other is active. | Hering et al. [14] |
Table 2: Continued.

| Global Geochemistry Mesh Impactor | Mesh Impactor | Polypropylene mesh | 2.4 | 1.5–1.7 | Hering et al. [14], Krupa [15] |
|----------------------------------|---------------|--------------------|-----|---------|-----------------------------|
| Sampler with cooling chamber     | Stainless steel cooling chamber | 5–30/30–120 | 0.2 | The volume of the sample collected depends on the duration of the fog event. | Hering et al. [14], Krupa [15] |
| Passive collector                | Nylon strings, 0.2 mm diameter | Collection area | 314 m² | Lange et al. [19] |
| AMC/WPI                          | Teflon strands | Baffle system. | Daube et al. [20] |
| Passive sampling system          | Teflon strings, 0.3 mm diameter | 15–25/120 | The sampler is activated and closed by a fog sensor based on the dew point and a separate rain sensor. | Krupa [15] |
| Automated sampling system        | Teflon strings | The measurement of pH and conductance occurs automatically in real time during sampling. | | Baumgardner et al. [3], Mohnen et al. [42], Vong et al. [43] |
| CRAC                             | 2 | The instrumentation and the electronic section include a conductometer, a pH meter and a microprocessor. | Jacob et al. [22] |

9. AUTOMATIC DEVICE

Figure 16 provides an example of a passive sampling system [15]. It consists of a Teflon support structure and 0.3 mm diameter Teflon strings, mounted 3.0 mm apart in a cylindrical configuration. Under appropriate airflow conditions, fog droplets are impacted on these strings, grow to larger drops, run down the strings, and are collected into bottles. All droplets > 5.0 µm diameter are impacted at normal wind speeds. In order to collect 15–25 mL it is necessary to collect samples for two hours. This collector is set out only in case of the occurrence of fog. In other situations it is enclosed inside a metal cylinder in order to prevent its contamination (rain and dry deposition). The sampler is activated and closed by a fog sensor based on the dew point and a separate rain sensor. A modified dynamic version of the passive sampler is the CalTech Active Strand Cloud Water Collector.

An automated system for the collection of cloud water samples directly from clouds (Figure 17) has been described in various papers [3, 42, 43]. The system consists of a collector which uses wind speed to affect cloud impact on 0.4 mm Teflon strings, a system for collecting, retaining, and storing samples, and an electronic unit controlling the system. The equipment includes also a temperature sensor, a rain detector, and a device for measuring wind speed. When there are no clouds, it is stored within a protective enclosure. During cloud events, a motor-driven shaft elevates and exposes the collector. The sample storage unit
Figure 15: Structure of a device for electrostatic precipitator for sampling single cloud or fog water droplets [15].

Figure 16: Structure of a passive fog water sampling system [15].

consists of 24 l polyethylene sample bottles contained in a circular wire support and housed in a commercial refrigerator. When the liquid water content of a cloud exceeds 0.05 g/m³, the wind speed is higher than 2.5 m/s, ambient air temperature is above freezing, and there is no rainfall, the cloud water collector is activated and projected out of its protective housing. The measurement of pH and conductance occurs automatically in real time during sampling. The indications of pH values are checked on a daily basis and the conductance value is indicated in relation to the external temperature.

An automated system for collecting and analyzing rain samples (Cloud and Rain Acidity/Conductivity Analyzer (CRAC)) has been adapted for the analysis of cloud water [5]. The system consists of a rain probe connected to an active CalTech collector [22] (Figure 18). When cloud water samples are being collected, the rain detector is disconnected. The instrumentation and the electronic section include a conductometer, a pH meter, and a microprocessor. Samples are collected sequentially in double accumulation vessels containing conductometric cells. After having collected 50 mL of samples, the system directs 12 mL to a chamber where pH is measured, while the remaining quantity goes to a vial which is kept in an automatic whirling arm. The whirling arm and the pH meter are situated in the cooling section. This collector has a 50% cutoff size centered around 2 µm diameter.

In published papers [47, 48], an automatic device for collecting fog samples has been described. Figure 19 shows a block diagram of this instrument [48, 49]. The microprocessor, after receiving signals from three sensors: fog, temperature, and rain, uses them to control the system of fog droplet collection. The fog detector, being an optical backscattered sensor (Figure 20), consists of two receivers; one of which measures the intensity of the light source and the other the intensity of the light scattered by the fog droplets. Both signals reach the microprocessor and their ratio is compared with a threshold value. The system is activated when the detector signals the presence of fog. An active string collector was used to collect fog samples because of its simple structure and ease of automation. This collector consists of a polyethylene aerodynamic tunnel in which air is sucked in by
Dew typically forms on cool nights with light breezes; water condenses from the atmosphere at ground level under these conditions. Hence, ground effects control the atmospheric chemistry during condensation [51]. Dew can play an important role in the deposition of air pollution particularly in arid ecosystems [18]. Dew may increase seedling survival, plant growth, and crop yield, but it may also have a negative effect, promoting bacteria and fungal infections. As a source of information on the environment, dew samples have long been a subject of study [52]. During the night, the latent heat flux towards the soil surface is very small, and therefore the amounts of dew deposition are very small as well. This fact poses special technical measurement difficulties that we will discuss in the following paragraphs.

The first publications presenting methods for determining the amount of dew appeared at the end of the last century [52]. Measurements were carried out using very simple methods, that is, collecting dew from grass by means of a sponge or by placing absorbent paper on grass. The dew samples were collected early in the morning. Before the expected appearance of dew, the collecting surface was flushed with deionized water and subsequently dried. Dew collection took place only on rainless nights to eliminate any influence of rain droplets on collected dew samples.

Modern methods of dew collection can be divided into three basic groups: optical, volumetric, and gravimetric. Modern methods permit continuous registration of dew events in their nascent stage, measurement of amount of dew, its duration, and evaporation time. Optical methods consist in visually estimating the amount of dew. In this method the device Duvdevan is often used.

Volumetric methods are connected with measuring the amount (volume) of the collected dew. Volumetric methods are exemplified by the volumetric drosometer, which has a collecting surface consisting of a filter paper 9 cm in diameter, saturated with water [53]. This instrument has limitations: it can be used only at temperatures above 0°C and the exact moment of dew formation must be determined.

Gravimetric methods rely on defining the increase of the weight of the collecting surface (without and with dew), which can be determined by means of analytical balance. Gravimetric methods include the method using Leick’s plates (made from a mixture of silicon dioxide dust, alabaster gypsum, and water) [53]. The plates are weighed before being used and after they are bedewed. The increase in weight in mg is proportional to the amount of dew in mm. Gravimetric methods include the drosograph too. The nascent dew settles upon a receiving plate and then flows down to a collecting vessel located underneath. The plate and vessel are located at one end of a first-order lever, the opposite end of which ends in a small writing pen. The movement of the pen, combined with the rotating movement of the drum, makes it possible to record changes in the quantity of dew. The drosograph described by Hutorowicz [54] uses an analytical balance with a metal box with turf in the role of dew collector and holder.

Li [55] has described another direct gravimetric technique. The collectors are containers made of polymethylmethacrylate (PMMA), with a diameter of 9.2 cm and a height of 10 cm, filled with an appropriate collecting material: thick gravel, sand, and/or loess.

As with cloud and fog, dew can deposit atmospheric pollution and in recent years, interest has grown in dew sampling for analysis of atmospheric pollution. The dew sampling method most often used for experiments on atmospheric pollution is the cloth plate method (CPM) [18, 56], a variant of the gravimetric method of dew collection. Velvet-like fabric (a square cloth of 6 cm × 6 cm), 0.15 cm thick, is placed in the centre of a 10 cm × 10 cm × 0.2 cm glass plate.
The glass plate is placed on a 10 cm × 10 cm × 0.5 cm layer of plywood. The plate and the plywood form a homogenous base 0.7 cm thick. The absorbent material is removed and replaced every day. Upon removal it is placed in flasks and weighed in order to measure dampness.

In the collector described in paper [57] and shown in Figure 21, the condensation surface is a rectangular foil sheet, 3 m × 10 m, made from TiO₂ and BaSO₄ microspheres embedded in polyethylene (volumetric method). The foil is fixed by lateral cables on a light grid attached by cables. The cables are fixed to beams anchored to the ground. This foil bedded in polyethylene (volumetric method). The foil is exhibited improved emitting properties in the near infrared (to provide radiative cooling of room temperature surfaces) and efficiently reflects the visible radiation (sun). A weak wind (< 1 m/s) is necessary to provide sufficient humid air around the condenser, but strong wind increases heat losses. To minimize wind influence and recover water drops by gravity using a plane condensing area with an angle with respect to horizontal and thermally isolated from the ground with 2 cm thick polystyrene foam, the placement of the collector at an angle of 30 degrees facilitates the flow of droplets (it seems that the angle may be a critical value, too high leads to diminishing dew formation). Dew, accumulating in a groove along the lower edge of the collecting plane, flows off to a 25 l polyethylene bottle.

Dew has been sampled by means of a collector which consists of a pump and a Teflon pipe terminated with a glass fibre filter [64]. The dew is sucked in by the pump together with air and flows through the pipe into a polyethylene bottle. Within 30 minutes, 1 mL of dew is collected.

In Table 3 parameters of samplers used for collecting dew were set together.

11. COLLECTING HOARFROST/RIME SAMPLES

Rime and hoarfrost are physical processes and forms when supercooled cloud droplets in a basal cloud layer freeze on impact with vegetation or topographic surfaces. Rime is commonly observed on trees, towers, power lines, and other objects at high elevations exposed to high-velocity cloud airflow. The difference between rime and hoarfrost was explained in Table 1. The chemical characteristics of rime, hoarfrost, and glaze should mimic the chemistry of cloud droplets and potentially affect vegetation adversely. If these deposits act as a biologically inert, frozen protective shell, their role could be beneficial. Hoarfrost and rime constitute an important element in water circulation, particularly in mountainous regions where they occur quite frequently, they contribute to the process of cleaning the atmosphere and of transferring impurities from air to the soil. Such collectors can be used interchangeably because the precipitation depends on meteorological conditions.

The simplest samplers used to collect samples of hoarfrost and rime consist of flat surfaces made from materials such as Teflon, galvanized steel [79], or Nylon wires [80]. Very often hoarfrost accumulates on some elements of cloud water collectors, for example, the strings, and it is then collected manually into polyethylene bottles [30, 81]. In Table 4 parameters of samplers used for collecting rime were set together.

In another published paper [60] an apparatus was described as being consisting of four polyethylene plates. The plates are fastened at the height of 1 m on an aluminium scaffold and positioned vertically facing the four cardinal points. The wind direction is estimated during sample collection by observing the wind vanes. Hoarfrost or else rime which accumulates on the surface of the plates is removed by means of a polyethylene scraper from the outer sides of the plates and collected into bottles.

For collecting samples of hoarfrost and rime, a passive collector is used with a string polyethylene screen. The collector mesh has 12 mm openings with 2 mm strands. Samples gathering on the screen are collected into 250 mL containers [61].

In the passive shield hoarfrost/rime collector is made of polyvinyl chloride and the collecting area consists of 46-singular fibres 0.2 mm in diameter. Deposit is scraped off from its surface by means of a scraper. Subsequently, the samples are placed in polyethylene bottles and transported to the laboratory [62].

To measure liquid and solid atmospheric deposits, the instrument known as the Grunow thimble is used also. It consists of a wire mesh in the form of a cylinder with a diameter of 10 cm and a height of 20 cm [82, 83]. This thimble is laid over a rain gauge, for example, of the Hellman type (the instrument consists of a metal cylinder ending in a funnel and the precipitation—depending on its kind—flows down into the container or accumulates above the funnel).

12. AUTOMATIC DEVICE FOR SOLID DEPOSITS

Measurements of solidified atmospheric deposits (hoarfrost, rime, freezing rain, and glazed frost) can be of substantial significance for certain branches of economy, such as the power industry, or air and road transport. It must be said that some problems may arise to differentiate between solid deposited precipitation and freezing fallen precipitation (freezing rain), both contributing to the icing events. A sufficiently thick layer of the deposit can overload and possibly break electric power lines. In a published paper [84] a method is presented for measuring the process of accretion of solid deposits on specific surfaces. The method relies on the measurement of the weight of ice deposits, the visual description of their appearance, and the time and duration of their occurrence. For this purpose, several pairs of electric power
### Table 3: Basic parameters characterizing dew collectors.

| Type of collector surface | Dimension (mm × mm) | Thickness of collector base (mm) | Comments | Reference |
|---------------------------|---------------------|----------------------------------|----------|-----------|
| PTFE-sheet                | 310 × 307           | 3                                |          |           |
| Pyrex glass plate         | 300 × 298           | 3                                |          |           |
| Stainless steel sheet     | 303 × 228           | 0.5                              |          | Takenaka et al. [58] |
| Aluminium sheet           | 303 × 226           | 1                                |          |           |
| PTFE-coated stainless steel sheet | 303 × 228 | 0.5                              |          |           |
| PTFE-coated aluminium sheet | 303 × 226 | 1                                |          |           |
| Plastic foil              | 2000 × 2000         | 0.05                             | Dew drops form on surfaces slightly inclined down the foil, they are collected in the morning by means of a syringe and put into 100 and 250 mL bottles. | Scheller [59] |
| Glass                     | 1000 × 1000         | 1                                | In order to insure that dew will form, the back of the collector is made of aluminium. | Jiries [51] |

### Table 4: Basic parameters characterizing RIME collectors.

| Type of collector surface | Dimension (cm²) | Location | Comments | Reference |
|---------------------------|-----------------|----------|----------|-----------|
| Polyethylene plates       | 400             | At the height of 1 m | It is also possible to measure the size of the collected deposit and to record its visual appearance by photographing it. | Ferrier et al. [60] |
| Polyethylene screen       | 3600            | At the height of 2–3 m | Deposits are collected for the whole 24-hour period. | Berg et al. [61] |
| Polyvinyl chloride shield | 929             | At the height of 1 m | Teflon film attached to a 3 cm thick polystyrene block with double-stick tape. | Duncan [62] |

#### References
- Takenaka et al. [58]
- Scheller [59]
- Jiries [51]
- Ferrier et al. [60]
- Berg et al. [61]
- Duncan [62]
- Foster [63]

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### Table 5: Information on the concentrations of inorganic and organic pollutions in fog, cloud, dew, and rime samples.

| Sampler | Atmospheric precipitation or deposits | Analytes | Concentration range (mg/L) | Reference |
|---------|--------------------------------------|----------|---------------------------|-----------|
| CalTech Rotating Arm Collector | Fog | NO$_3^-$ | 0.37–489.8 | Jacob et al. [41] |
| | | SO$_4^{2-}$ | 2.59–99.84 | Johnson et al. [40] |
| | | Cl$^-$ | 0.67–194.9 | Lacaux et al. [65] |
| | | NH$_4^+$ | 0.77–51.48 | |
| | | Ca$^{2+}$ | 0.6–39.0 | |
| | | K$^+$ | 0.08–5.77 | |
| | | Mg$^{2+}$ | 0.11–41.25 | |
| | | Na$^+$ | 0.23–139.4 | |
| | | pH | 2.16–6.17 | |
| | | Formaldehyde | 0.007–0.17 | Grosjean and Wright [66] |
| | | Acetaldehyde | 0.08–0.32 | |
| | | Benzaldehyde | 9.98–1010.6 | Waldman et al. [12] |
| | | NO$_3^-$ | 6.14–446.4 | Grosjean and Wright [66] |
| | | SO$_4^{2-}$ | 0.53–343.1 | |
| | | Cl$^-$ | 1.11–133.6 | |
| | | NH$_4^+$ | 0.02–6.88 | |
| | | Fe | 0.038–2.78 | |
| | | Pb | 2.06–3.87 | |
| | | pH | 0.01–1.08 | |
| | | Formaldehyde | 0.007–0.17 | Grosjean and Wright [66] |
| | | Acetaldehyde | 0.08–0.32 | |
| | | Benzaldehyde | 9.98–1010.6 | Waldman et al. [12] |
| | | NO$_3^-$ | 6.14–446.4 | Grosjean and Wright [66] |
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| | | Acetaldehyde | 0.08–0.32 | |
| | | Benzaldehyde | 9.98–1010.6 | Waldman et al. [12] |
| | | NO$_3^-$ | 6.14–446.4 | Grosjean and Wright [66] |
| | | SO$_4^{2-}$ | 0.53–343.1 | |
| | | Cl$^-$ | 1.11–133.6 | |
| | | NH$_4^+$ | 0.02–6.88 | |
| | | Fe | 0.038–2.78 | |
| | | Pb | 2.06–3.87 | |
| | | pH | 0.01–1.08 | |
| CASCC | Fog | NO$_3^-$ | 4.72–1791.8 | Munger et al. [32] |
| | | SO$_4^{2-}$ | 0.013–1196.16 | Klemm et al. [67, 68] |
| | | Cl$^-$ | 0.0027–86.51 | Collett et al. [69] |
| | | NH$_4^+$ | 0.38–464.4 | Anastasio et al. [70] |
| | | Na$^+$ | 0.21–44.48 | Wrzesinsky et al. [71] |
| | | K$^+$ | 0.23–2.67 | Herckes et al. [35, 72] |
| | | Al | 14–903 µg/L | |
| | | Ni | 0.8–42.8 | |
| | | Pb | 3.4–61.4 | |
| | | Cd | 0.3–8.7 | |
| | | Cu | 2.5–56.8 | |
| | | Sb | 0.8–5.7 | |
| | | Se | 1.1–11.5 | |
| | | pH | 2.33–7.43 | |
| | | Conductivity | 17–452 µS/cm | |
| | | n-Alkanes | NW-5.1 ng/mL | |
| | | Naphthalene | NW-0.1 | |
| | | Phenanthrene | 0.028–0.169 | |
| | | Anthracene | NW-0.040 | |
| | | Fluoranthrene | 0.002–0.095 | |
| | | Pyrene | 0.013–0.289 | |
| | | Benzo(ghi)fluoranthene | NW-0.019 | |
| | | Benz(a)anthracene | NW-0.022 | |
| | | Benzo(e)pyrene | NW-0.252 | |
| | | Benzo(a)pyrene | NW-0.218 | |
| | | Indeno(1,2,3-cd)pyrene | NW-0.007 | |
| | | Benzo(ghi)perylene | NW-0.033 | |
### Table 5: Continued.

| Concentrations | Units | References |
|---------------|-------|------------|
| Cloud NO$_3^-$ | 1.74–507.8 | Munger et al. [34] |
| SO$_4^{2-}$    | 36.48–1112.8 | Anastasio et al. [70] |
| Cl$^-$         | 0.39–74.4 | Collett et al. [73] |
| NH$_4^+$       | 0.79–168.9 | Igawa et al. [33] |
| Ca$_2^+$       | 0.24–91.4 |            |
| Mg$_2^+$       | 0.11–126.6 |            |
| Na$^+$         | 0.32–87.3 |            |
| pH             | 2.42–4.98 |            |
| Formaldehyde   | 0.41–1.84 |            |
| Acetaldehyde   | 0.053–0.22 |            |
| ASRC Fog NO$_3^-$ | 0.22–23.15 | Schemenauer et al. [74] and Schemenauer and Cereceda [75] |
| SO$_4^{2-}$    | < 0.5–36.4 |            |
| F$^-$          | < 0.1–1.48 |            |
| Cl$^-$         | < 0.1–27.9 |            |
| Ca$_2^+$       | < 0.10–11.10 | Gordon et al. [9] |
| K$^+$          | < 0.08–0.93 | Eckardt et al. [76] |
| Mg$_2^+$       | 0.02–1.79 |            |
| Na$^+$         | 0.08–15.7 |            |
| NH$_4^+$       | 0.32–3.19 |            |
| Al             | 0.02–1.56 |            |
| B              | < 0.07–0.22 |            |
| Cd             | < 0.002 |            |
| Cr             | < 0.01 |            |
| Cu             | < 0.01 |            |
| Mn             | < 0.003–0.236 |            |
| Ni             | < 0.01 |            |
| Pb             | < 0.03–0.27 |            |
| Zn             | < 0.01–1.03 |            |
| HCO$_3^-$      | 0.31–1.86 |            |
| pH             | 3.46–6.694 |            |
| NO$_3^-$       | 0.70–12.3 | Schemenauer and Cereceda [77] |
| SO$_4^{2-}$    | 0.45–6.70 |            |
| Cl$^-$         | 5.85–84.3 |            |
| NH$_4^+$       | 0.00–0.65 |            |
| Ca$_2^+$       | 2.08–42.7 |            |
| K$^+$          | 0.14–2.12 |            |
| Mg$_2^+$       | 0.46–5.73 |            |
| Na$^+$         | 3.36–46.0 |            |
| Fe             | < 0.06 |            |
| Cd             | < 0.0005 |            |
| Pb             | < 0.0005 |            |
| Be             | < 0.0005 |            |
| Cr             | < 0.005 |            |
| Mn             | 0.0054–0.034 |            |
| Ni             | < 0.002–0.0074 |            |
| Cu             | < 0.005 |            |
| Zn             | < 0.002–0.026 |            |
| pH             | 6.96–7.94 |            |
| High-Volume    | Diazinon 0.15–4.8 µg/L | Schomburg et al. [27] |
| Sampler        | Malathion 0.14–8.7 | Chernyak et al. [28] |
|                | Chlorpyrifos 0.012–0.19 |            |
|                | Fonofos < 0.012–0.030 |            |
|                | Methidathion 0.036–0.22 |            |
|                | Chlorpyrifos < 900–5000 |            |
| Table 5: Continued. | Chlorothalonil | 4000–17000 |
|---------------------|----------------|-----------|
|                     | Endosulphan    | 500–10000 |
|                     | Metolachlor    | 1800–147000 |
|                     | Trifluralin    | 100–1000   |
| NO$_3^-$            | 0.248–284.0    | Vermeulen et al. [29] |
| SO$_4^{2-}$         | 0.528–163.2    | Elias et al. [30] |
| F$^-$               | 0.02–0.67      | Kimball et al. [31] |
| Cl$^-$              | 0.60–24.7      |            |
| Ca$^{2+}$           | 0.02–16.0      |            |
| Na$^+$              | 0.09–8.33      |            |
| K$^+$               | 0.12–19.5      |            |
| Mg$^{2+}$           | 0.06–5.5       |            |
| NH$_4^+$            | 0.054–80.5     |            |
| Al$^{3+}$           | 0.04–1.16      |            |
| Li$^{3+}$           | 1.00–2.00      |            |
| Sr$^{2+}$           | 0.03–0.05      |            |
| pH                  | 2.41–6.36      |            |
| Conductivity        | 23.2–854 µS/cm  |            |

| CWP Fog             | SO$_2^{+}$     | 0.88      |
|                     | Cl$^-$         | 0.65      |
|                     | Na$^+$         | 0.29      |
|                     | K$^+$          | 0.14      |
|                     | Mg$^{2+}$      | 0.06      |
| Rime                | SO$_4^{2-}$    | 0.31      |
|                     | NO$_3^-$       | 0.77      |
|                     | pH             | 5.69      |
|                     | Conductivity   | 6.65–252 µS/cm |

| Passive collector   | SO$_4^{2-}$    | 0.88      |
| (polyvinyl chloride)| Cl$^-$         | 0.65      |
|                     | Na$^+$         | 0.29      |
|                     | K$^+$          | 0.14      |
|                     | Mg$^{2+}$      | 0.06      |

| CPM Dew polyethylene| Ca$^{2+}$      | 0.21      |
|                     | NH$_4^+$       | 0.31      |
|                     | NO$_3^-$       | 0.77      |
|                     | pH             | 5.69      |
|                     | Conductivity   | 6.65–252 µS/cm |

| CPM Teflon Dew      | NO$_3^-$       | 2.16–8.14 |
|                     | NO$_3^-$       | 5.77–14.76 |
|                     | SO$_4^{2-}$    | 8.5–43.25 |
|                     | Cl$^-$         | 0.99–3.05 |
|                     | NH$_4^+$       | 5.68–14.06 |
the environment. Once every minute a signal appears at the output of the sensor indicating the frequency of vibrations of the sensor.

13. MEASUREMENT RESULTS

In the recent years an increased interest is observed in the chemistry of atmospheric precipitation and deposits, as the impurities and pollutants undergo complicated chemical and biochemical reactions in the aquatic and soil ecosystems due to which they enter into biogeochemical circulation, disturbing the environmental balance. For this reason, the pollution of atmospheric air, as well as the pollution of atmospheric precipitation and deposits which follow, constitutes a problem on an international scale, requiring constant monitoring as being confirmed by national and foreign studies collected through literature research. In Table 5 results of measurements of inorganic and organic compound concentrations in nontypical samples are presented.

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Table 5: Continued.

|    |     |
|----|-----|
| Ca²⁺ | 7.54–13.46 |
| K⁺  | 0.27–3.12  |
| Mg²⁺ | 0.69–2.83  |
| Na⁺ | 0.34–2.94  |
| pH  | 5.4–6.6   |
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