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Streaming potential in volcanic rocks from Mount Pelée

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Abstract. Streaming potential and electric conductivity have been measured in a laboratory on 11 consolidated samples coming from five deposits of the different evolutionary stages of Mount Pelée volcano. The streaming potential coupling coefficient ranges from -35 to -4905 mV MPa⁻¹ and increases with increasing permeability. This increase is mainly due to the dependency of rock effective conductivity with permeability. The permeability of the samples varies from 0.146×10^{-12} to 34×10^{-12} m². The zeta potential, at pH = 7 and water conductivity of 2.1×10^{-4} S m⁻¹, is relatively small for the majority of the samples. It ranges from -4 to -19 mV. According to water conductivity analysis on Mount Pelée, streaming potential coupling coefficients of -25 to -406 mV MPa⁻¹ can be expected for this volcano.

1. Introduction

Monitoring of electric and magnetic anomalies has been proposed as a means of predicting earthquakes [Mizutani et al., 1976; Fenoglio et al., 1995; Trique et al., 1999] or volcanic eruptions [Zlotnicki et al., 1998]. Magnetic observations made on La Soufrière volcano (Guadeloupe, France) have shown magnetic variations that could be induced by electric fields related to deepwater circulation (approximately at a depth of 5 km) [Pozzi et al., 1979]. Water circulation is also thought to be the main agent of the strong magnetic signals observed on Piton de La Fournaise volcano (Réunion Island, France) [Zlotnicki and Le Mouël, 1990].

Self-potential (SP) anomalies can be produced by a streaming potential, as a consequence of fluid pressure gradients (an electrokinetic phenomenon); by a thermoelectric potential, as a consequence of a temperature gradient; by a chemical potential, as a consequence of a chemical gradient; or by these effects possibly acting together. Indeed, SP anomalies have been shown to be related to hydrothermal activity (two-phase water and steam flow) in geothermal areas and on active volcanoes [Zohdy et al., 1973; Anderson and Johnson, 1976; Corwin and Hoover, 1979; Michel and Zlotnicki, 1998].

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Electrokinetic phenomena were proposed as a possible mechanism for generating SP anomalies observed in geothermal areas [Corwin and Hoover, 1979; Revil and Pezard, 1998]. Some significant SP variations have been observed associated with volcanic activity and can precede eruptive episodes [Aubert and Kieffer, 1984; Malengreau et al., 1994]. SP anomalies are commonly thought to be induced by electrokinetic phenomena because, thermoelectric and chemical coefficients are smaller than the electrokinetic coefficient [Nourbehecht, 1963; Sill, 1983; Fitterman, 1979; Perrier et al., 1998, 1999]. SP surveys performed on Etna. Lamonga, and Merapi volcanoes have shown that the amplitude of the anomalies was related to the vapor flow, and the main source of these SP anomalies was thought to be electrokinetic [Aubert and Dana, 1994]. In some areas there is a good spatial correlation on active volcanoes between rising two-phase convective cells and positive SP anomalies [Aubert et al., 1984; Aubert and Baubron, 1988]. SP measurements on La Fournaise volcano (Réunion Island, France) have shown a huge positive anomaly of amplitude more than +1850mV on the cone. This anomaly suggests that hydrothermal ascendant circulations within the cone are induced by a magmatic complex source at least at 2 km depth beneath the summit [Zlotnicki et al., 1994]. Recent SP measurements on Unzen volcano (Kyushu Island, Japan) showed a positive anomaly as large as +1000 mV per 500 m in the vicinity of the newly extruded lava dome. The most reasonable mechanism

for this anomaly is thought to be streaming potentials associated with subsurface hydrothermal convection [Hashimoto and Tanaka, 1995]. A sharp increase of +500 to +600 mV in SP was detected 3 months preceding the first extrusion of lava and was considered to be a result of the growth of the hydrothermal system. Moreover, Fujinawa et al. [1992] measured an anomalous vertical electric field of about +30 mV a few days prior to, and a month after, a minor volcanic eruption of Mount Mihara. These anomalous variations were thought to be generated by electrokinetic phenomena induced by variations of hydrothermal circulation around the crater, or by variations of crack density, due to volcanic activity.

Modeling of all these observations needs a good understanding of electrokinetic phenomena in volcanic rocks, specially the effect of the most influential parameters on streaming potential. These effects can be quantified in the laboratory. Few laboratory data of geophysical interest on streaming potential are available [Ahmad, 1964; Somasundaran and Kulkarni, 1973; Ishido and Mizutani, 1981; Jouniaux et al., 1994; Pozzi and Jouniaux, 1994; Jouniaux and Pozzi, 1995a, b, 1997; Lorne et al., 1999a, b], and few can be applied directly to volcanic environments [Massenet, 1983; Massenet and Van Ngoc, 1985; Antraygues and Aubert, 1993].

We present in this paper laboratory measurements of streaming potential and electric conductivity on 11 volcanic samples representative of the main lithologies and textures of the Mount Pelée volcano.

2. Electrokinetic Phenomena

Electrokinetic phenomena are induced by the relative motion between the fluid and the rock. Minerals forming the rock develop an electric double layer when in contact with an electrolyte, usually resulting from a negatively charged mineral surface. An electric field is created perpendicular to the surface of the mineral which attracts counterions (usually cations) and repulses anions in the vicinity of the pore-matrix interface. The electric double layer is made up of the Stern layer, where cations are adsorbed on the surface, and the Gouy diffuse layer, where the number of counterions exceeds the number of anions (for a detailed description, see Adamson [1976] and Hunter [1981]). The streaming potential is due to the motion of the diffuse layer induced by a fluid pressure difference along the interface. The zeta potential is defined at the slipping plane or shear plane (i.e., the potential within the double layer at the zero-velocity surface). In a porous medium the electric current density and the fluid flux are coupled [Overbeek, 1952; Nourbehecht, 1963], so that the streaming potentials are generated by fluids moving through porous media.

The parameter that quantifies this coupling is the streaming potential coupling coefficient or, simply, the coupling coefficient, defined by

$$C_S = \frac{\Delta V}{\Delta P} = \frac{\epsilon \zeta}{\eta \,\sigma_{\text{eff}}} \,, \tag{1}$$

where ΔV is the generated electrical potential, ΔP is the applied pore pressure difference, ϵ is the dielectric constant of the pore fluid ($\epsilon = 7 \times 10^{-10}$ F m⁻¹), ζ is the zeta potential, η is the dynamic viscosity of the pore fluid ($\eta = 10^{-3}$ Pa s), and $\sigma_{\rm eff}$ is the effective conductivity defined by

$$\sigma_{\rm eff} = F \,\sigma_r \,, \tag{2}$$

where F is the formation factor and σ_r is the rock electrical conductivity at the fluid conductivity used for electrokinetic measurements.

If surface conductivity is negligible, $\sigma_{\rm eff} = \sigma_f$, and we have

$$C_S = \frac{\epsilon \zeta}{\eta \, \sigma_f} \,, \tag{3}$$

which is the Helmholtz-Smoluchowski equation [Ishido and Mizutani, 1981]. When we want to be free from the effect of the effective conductivity, another parameter, C', independent on this conductivity is defined,

$$C' = \frac{\Delta V}{\Delta P} \,\sigma_{\rm eff} = \frac{\epsilon \,\zeta}{\eta} \,, \tag{4}$$

where C' is called the electrokinetic coefficient.

For a complete development of the equations governing the coupled electro-magnetics and flow of the porous media, see *Pride* [1994] and *Revil et al.* [1999a, b], for modeling of electrokinetics in porous rocks see *Coelho et al.* [1996] and *Bernabé* [1998], and for further details on surface conductivity, see *Revil and Glover* [1997, 1998].

3. Experimental Procedures

Streaming potential and electric conductivity measurements have been performed on 11 consolidated samples coming from five deposits of the different evolutionary stages of Mount Pelée. This explosive volcano, located in the north end of Martinique Island in the Lesser Antilles (France), is mainly composed of andesitic pyroclastic materials. The sampling represented different lithologies and textures typically found on this volcano (dense or porous block, pumice, and scoria). Mineralogical and chemical compositions are relatively constant [Westercamp, 1976; Fichaut et al., 1989; Vincent et al., 1989]. Plagioclase, orthopyroxene, clinopyroxene, and a few magnetite are the main phenocrystals (with the following mean proportion: 33% plagioclase, 8% pyroxene, and 2.8% magnetite). The groundmass of the samples ranges from 52 to 63%. It is composed of variable percentages of glass and microcrystals. The proportion of the different mineral phases and the groundmass varies from one eruption to another. A short description of the samples is presented in Table 1.

3.1. Sample Preparation

Two samples were cored on each block of rock: (1) cylinder, 23 mm in diameter and 25–50 mm length, for porosity, permeability, and electrical conductivity measurements, and (2) cylinder, 2.5 cm in diameter and 48 mm long, for streaming potential measurements. The

Stage	Eruption	Sample	Deposit	Description
Paleo-Pelée		MA701 (1) MA701 (3)	volcanic breccias	vesicular andesite blocks
Neo-Pelée	Saint Vincent	MF201 (1)	scoria flows	andesite scoriaceous blocks
	Saint Vincent	MF201 (2)	scoria flows	andesite scoriaceous blocks
	Saint Vincent	MF201 (3)	scoria flows	andesite scoriaceous blocks
Modern stage	P1	F3	Pelean nuees ardentes	lightly vesicular andesite block
	P1	M	Plinian fallout	andesitic pumice
	1902	K	Pelean nuees ardentes	vesicular grey andesite
	1902	J	Plinian fallout	white andesitic pumice
	1929	R	Block-and-ash flows	vesiculated andesite
	1929	Q	Block-and-ash flows	highly vesicular andesite

 Table 1. Description of the Samples

end faces of the samples were ground flat and parallel to within 0.025 mm.

3.2. Porosity and Air Permeability Measurements

The matrix mass density ρ_m was measured with a pycnometer on 2-3 g of rock reduced on fine powder (10 μ m). The porosity was measured with the triple weighing method: the sample was first oven dried at 70°C for at least 48 hours, slowly cooled to room temperature, and then saturated with degassed water under vacuum. The sample is weighed before saturation, and after saturation and immersed (saturated sample) within the water. These three measurements and the matrix mass density allow us to calculate the connected, ϕ_C , and trapped porosity, ϕ_T . The trapped porosity is the part of porosity not accessible under vacuum. The experimental error on porosity was less than 1%.

The air permeability k was measured, at atmospheric pressure, with a falling head permeameter (see *Bourbié* and Zinszner [1985] for a description of the apparatus). The accuracy is within 1% for permeabilities greater than 10^{-16} m². The Klinkenberg effect is only significant in the three less permeable samples, and it is always less than 6%. Measurements were repeated several times for reliability.

3.3. Electric Conductivity Measurements

In order to determine the electrical formation factor F, the electrical conductivity of the rock was measured, at room conditions, when the samples were saturated successively with a distilled water and with at least seven aqueous NaCl solutions with conductivities increasing from 6×10^{-4} to 17 S m⁻¹. The electrical conductivity measurements were performed with twopole electrodes, a technique consisting in the measure of the electrical impedance of the samples, at different frequencies (varying from 100 to 100 kHz), by placing the sample between two stainless steel electrodes, connected to an impedance meter (HP4263A). Filter paper disks soaked with the saturating solution were put between the electrodes and each end of the sample to reduce contact resistance and to prevent polarization effects. A clamping system held the sample between the electrodes with the same constant force for each sample. The electrolyte conductivity was measured with a conductivity cell (Knick 702).

The following experimental procedure was used:

1. The sample was first dried for at least 48 hours at 70°C, cooled to room temperature, and placed in a container under dynamic vacuum for 4 hours. The container was then filled with distilled and degassed water and left under vacuum for 48 hours.

2. Samples were inserted quickly inside an insulating adhesive jacket to limit water evaporation during measurements and to prevent electrolyte conduction on the external surface sample. The electrical impedance of the sample and the conductivity of the solution were then measured.

3. After these measurements the samples were dried and again saturated with a degassed NaCl solution of 2×10^{-4} mol L⁻¹, and the electrical measurements were performed. The saturation conditions and the procedure of measurements were those described in the previous paragraph.

4. This procedure was reiterated for the other six NaCl solutions. Between two measurements, with two different saline solutions, the samples were carefully cleaned by saturating them several times with distilled water. The relative experimental error was less than 1% and 0.5% for rock conductivity and fluid conductivity measurements, respectively.

The formation factor was determined by nonlinear inversion of the experimental data, using the *Revil and Glover* [1998] model:

$$\sigma_r = \frac{\sigma_f}{F} H(\xi) \,, \tag{5}$$

$$H(\xi) = 1 - t_{(+)}^{f} + F\xi + \frac{1}{2} \left(t_{(+)}^{f} - \xi \right) \\ \times \left(1 - \frac{\xi}{t_{(+)}^{f}} + \sqrt{\left(1 - \frac{\xi}{t_{(+)}^{f}} \right)^{2} + \frac{4F\xi}{t_{(+)}^{f}}} \right), \quad (6)$$

where $t_{(+)}^{f}$ is the Hittorf number of the cations (fraction

of the electrical current carried in the free electrolyte by the cations), F and Φ are the formation factor and the porosity, respectively, and the dimensionless parameter ξ is defined by *Kan and Sen* [1987]

$$\xi = \frac{\sigma_S}{\sigma_f} = \frac{2}{3} \left(\frac{\Phi}{1 - \Phi} \right) \frac{\beta_S Q_V}{\sigma_f} , \qquad (7)$$

where σ_S is the surface conductivity, β_S is the surface mobility of the counterions, and Q_V is the excess surface charges per unit of pore volume. The inversion procedure is described by *Bernard* [1999]. Only the data unaffected by the polarization effects were used for the inversion. For most of the samples the frequency which allows the electric impedance to be a pure resistance is 10 kHz.

3.4. Streaming Potential Measurements

The samples were first saturated with deionized water under vacuum for 8 hours. Electrical potential was measured while water was made to flow through the sample using the apparatus described in Figure 1. A silicone jacket was put around the sample to ensure the separation of the water and the confining fluid. A confining pressure (maximum 300 kPa) could be applied using the compressed air. The axial load was applied using lead weights of 50 kg, so that the axial stress on the sample was of the order of 1 MPa. This system allows a water to circulate through the sample with a maximum pressure gradient of 300 kPa. The water pressure, measured with a manometer, was controlled by applying compressed air pressure to a water reservoir at one end of the sample while the other end was maintained at atmospheric pressure. The tubing circuit was not closed, so that there was no possible electric current leakage through the water within the tubing. Most of the tubing is plastic, and the few metallic parts of the apparatus were grounded. The electric potential was measured by two silver-chloride electrodes of length 40 mm, which were made by electrolysis of a silver rod in salt water of concentration 10^{-1} mol L⁻¹. The electrodes were put in the water circuit near the ends of the sample but not within the water circulation to avoid the electrical noise due to water movement near the electrodes. The two electrodes are connected by two coaxial cables to a high-input impedance (above $10^{10} \Omega$) volt meter. The resistance of the sample was usually several hundred $k\Omega$ to a maximum of 1 M Ω , which is low compared to the input impedance of the volt meter, therefore allowing accurate measurements of the potentials.

Streaming potential measurements were performed once equilibrium in water pH and water conductivity was attained: first, deionized water was made to flow through the sample, then collected, transferred to the upstream reservoir, and flowed through the sample again. The same procedure was repeated until the conductivity and pH of the water after flowing through the sample were constant. The volume of water flowing through the sample during this procedure ranged between 5 and 1000 cm³, depending on the sample per-

water outlet silicon jacket confining sample pressure electrode water inlet

Figure 1. Apparatus used for electrokinetic measurements.

meability (volume of the sample is 23.6 cm³). Once the water conductivity and pH were constant, the streaming potential was measured for various water pressure gradients (10-180 kPa). The water expelled from the sample was also analyzed. Samples with permeability below 10^{-13} m² were not tested, because the water flow produced with the applied pore pressures would have

 Table 2. Connected and Trapped Porosities, Permeability, and Electrical Formation Factor for Each Sample

Sample	ϕ_C	ϕ_T	k	F	σ_r
R	32.8	3.1	10342	9	0.19
Q	38.0	3.8	34325	9	0.19
K	25.1	2.3	1604	17	0.22
MF201 (1)	36.3	2.6	1564	16	0.85
MA701 (3)	28.9	2.3	5124	19	0.41
J	57.5	6.8	393	31	0.68
F3	14.6	1.7	207	65	0.15
М	57.4	10.0	146	4 0	0.76
MF201 (3)	35.2	2.9	5765	9	1.34
MA701 (1)	24.0	0.1	257	18	1.01
MF201 (2)	28.0	1.6	1043	25	0.43

Note that ϕ_C indicates connected porosity, in percent, ϕ_T indicates trapped porosity, in percent, k indicates permeability, in 10^{-15} m², F indicates formation factor, and σ_r indicates the rock conductivity, in mS m⁻¹.

been too small to induce a measurable electrokinetic effect. For lower permeabilities an apparatus similar to the one used with the pulse method to measure low permeabilities is needed [Jouniaux et al., 1994].

4. Results and Discussion

4.1. Porosity, Permeability, and Electrical Conductivity

Connected porosity in the samples ranges from 14.6 to 57.5%, and trapped porosity can reach 10% (Table 2). Permeability ranges from 0.146×10^{-12} to 34×10^{-12} m². The most permeable samples have permeabilities higher than 10^{-11} m², which corresponds to very permeable materials such as sand.

Figure 2 shows an example of the dependence of the rock electrical conductivity versus the saturating fluid conductivity, with the fit obtained using the *Revil and Glover* [1998] model. Formation factor values are reported in Table 2 and range from 9 to 65. We note a tendency of F to decrease with increasing permeability as shown in Figure 3. The relationship between permeability and formation factor is detailed by *Bernard* [1999]. The rock conductivities σ_r , at the fluid conductivities used for electrokinetic measurements (obtained directly or by extrapolation using the Revil and Glover model) are inferior to 1 mS m⁻¹, except for the scoria MF201(3), where it reaches 1.3 mS m⁻¹ (Table 2).

4.2. Streaming Potential

The streaming potential has been measured for 4-11 differential fluid pressure, 3 or 4 times for each sample. Figure 4 shows only the last set of measurements (the three last sets of measurements giving the same results, the first set corresponding to the transitory phase). We see that the streaming potential is proportional to the driving pore pressure, as expected from (1). This linearity has been verified previously for sand [Ahmad, 1964,



Figure 2. Rock electrical conductivity versus water conductivity, for sample F3.

Lorne et al., 1999a] and sandstone samples [Jouniaux and Pozzi, 1997].

For each sample, the coupling coefficient C_S has been deduced from the mean (from the three or four sets of measurements) of the slopes of the straight lines streaming potential versus driving pore pressure, by linear fit to the data, using least squares regression. The values of these coupling coefficients (Table 3) range from -35 to -4905 mV MPa⁻¹ for a pH and conductivity of the water, in equilibrium with the the rock, varying from 6.2 to 6.9 and 0.23 to 2 mS m^{-1} , respectively.



Figure 3. Formation factor versus permeability for all the studied samples.



Figure 4. Measured streaming potential ΔV versus the applied pore pressure ΔP for six samples.

Figure 5 displays the variation of the coupling coefficient with porosity, formation factor, and permeability. No specific correlation is observed with porosity or formation factor. However, the coupling coefficient increases, by more than 2 orders of magnitude, with increasing permeability. This reliance of the coupling coefficient with permeability may be a consequence of a dependency of the effective conductivity $\sigma_{\rm eff}$ with permeability. Figure 6 exhibits this dependency. We observe a net decreasing of the effective conductivity, by more than 1 order of magnitude, with increasing permeability.

The dependency of the coupling coefficient on the permeability has been observed by Antrayques and Aubert [1993] on quartz sands (1.5 to 12 mm grain size) when these authors measured the electric potential along a vertical column of porous material in which upward vapor and a downward liquid water (vapor condensation) flows occurred, and by Jouniaux and Pozzi [1995a] on Fontainebleau sandstone samples, with permeabilities ranging from 0.15×10^{-15} to 1.2×10^{-12} m². Jouniaux and Pozzi [1995b] suggested that the decreasing coupling coefficient with the decreasing permeability of Fontainebleau sandstones could be induced by surface conductivity that may not be negligible, as first proposed for narrow cylindrical capillaries [Broz and Epstein, 1976]. In our samples, where the effective conductivity was measured instead of being inferred from surface conductance assumptions, we found a variation of the effective conductivity with permeability (Figure 6) of ~ 1.3 orders of magnitude. In order to be free

of the dependence on the effective conductivity, the parameter C', which is independent from effective conductivity, has been calculated using (4). This parameter varies from -0.4 to -8.5 mV(MPa Ω m)⁻¹ (Table 3). Figure 7 shows the C' parameter as a function of permeability. We observe that the electrokinetic coefficient still varies by a factor 20 in the studied samples, but no clear dependence with permeability is noted.

4.3. The Zeta Potential

The macroscopic zeta potential is deduced for each sample, using (1). The values obtained are given in Table 3. The zeta potential ranges from -0.6 to -12.2mV. These values are relatively low. Four samples have zeta potential, in absolute value, lower than 2 mV, indicating that for these samples the pH of the distilled water in equilibrium with these rocks is close to the pH_{pzc} . (Value pH_{pzc} represents the pH corresponding to the zero surface charge. At pH_{pzc} the zeta potential is zero). Massenet and Van Ngoc [1985] measured the zeta potential in samples of volcanic ashes (from Mount Etna). The mineralogical composition of these ashes is relatively close to the composition of our samples. They obtained values of zeta potential ranging from -26 to -34 mV for pH and fluid conductivity equivalent to these of our measurements. These values are superior to our values, but in the Mount Etna samples, $pH_{pzc} = 4.4$. These differences can emanate from differences in mineralogical composition and particularly from a different percentage of glass in the groundmass.

The zeta potential varies from more than 1 order of



Figure 5. Measured coupling coefficient C_S versus (a) permeability, (b) formation factor, and (c) porosity.



Figure 6. Measured effective conductivity σ_{eff} as a function of permeability.

magnitude in the studied samples. This variation cannot be due to the effective conductivity, since we take σ_{eff} into account to deduce the zeta potential. The effect of other parameters should be considered to explain this difference. The most influential parameters on the zeta potential are the mineralogical composition, the pH, the fluid conductivity, and the temperature [Li and De Bruyn, 1966; Ishido and Mizutani, 1981; Revil et al., 1999a].

All the measurements were performed at the same temperature (25°C). The pH of the deionized water, at the equilibrium with the samples, pH_e , ranges from 6.2 to 6.9 (Table 3). It depends on the rock mineralogy. Thus, for the samples of the 1929 eruption, $pH_e = 6.2$, while for Paleo-Pelée stage samples, $pH_e = 6.6$. From the 1902 and P1 eruptions, which present a succession of the Plinian and Pelean mechanisms, in the samples emanating from the Pelean mechanism, $pH_e = 6.4$. The two samples coming from the Plinian mechanism have different values of pH_e (6.7 and 6.9), problably because the Plinian samples exhibit more heterogeneous groundmass composition than the Pelean samples do (P. Besson, personal communication, 1999). The pH_e of the Saint Vincent stage samples are all different. This stage is characterized by variations of composition that are relatively important.

Water conductivity in equilibrium with the rock depends on the mineralogical composition and on the equilibrium pH and varies from 0.2 to 2 mS m⁻¹ (Table 3). In order to compare the ζ potential obtained on the different samples in the same physical conditions, we must calculate the values of this potential at the same values of pH and water conductivity (or water concentration).

In the case of quartz systems, at pH close to 7 and for solutions containing K^+ or Na^+ , the ζ potential may

Sample	σ_f	$\sigma_{ m eff}$	pH _e	C_S	C'	ζ	$\zeta_{ m cor}$
R	0.34	1.7	6.2	-4905 + 15	-8.5	-12.2	-19
Q	0.23	1.7	6.2	-1940 + 30	-3.3	-4.7	-11
K	0.23	3.7	6.4	-1078 + 4	-4.0	-5.7	-11
MF201 (1)	1.31	13.6	6.8	-454 + 40	-6.2	-8.8	-11
MA701 (3)	0.34	7.7	6.6	-335 + 28	-2.6	-3.7	-7
J	0.74	21.0	6.7	-193 + 50	-4.1	-5.8	-9
F3	0.69	9.5	6.4	-127 + 5	-1.2	-1.7	-7
М	2.00	30.3	6.9	-88+7	-2.7	-3.8	-5
MF201 (3)	0.54	12.1	6.6	-56+4	-0.7	-1.0	-4
MA701 (1)	0.78	18.1	6.6	-35+2	-0.6	-0.9	-4
MF201 (2)	0.46	10.7	6.5	-37+12	-0.4	-0.6	-5

Table 3. Fluid and Effective Conductivities, pH of the water, Coupling Coefficient, Electrokinetic Coefficient, and Zeta Potential for Each Sample.

Note that σ_f indicates conductivity of fluid, in mS m⁻¹, σ_{eff} indicates effective conductivity, in mS m⁻¹, pH_e indicates the pH of the water at the equilibrium with the rock, C_S indicates coupling coefficient, in mV MPa⁻¹, C' indicates electrokinetic coefficient, in millivolt (MPa Ω m)⁻¹, ζ indicates zeta potential from equation (1), and ζ_{cor} indicates zeta potential from equation (8), in mV.

be analytically expressed as a function of temperature, pH, and fluid concentration [Revil et al., 1999a] by

$$\zeta = c - d \,\mathrm{pH} + b \log_{10} C_f \,, \tag{8}$$

where C_f is the fluid concentration. The slopes b and d of (8) depend only on temperature. The zeta potential decreases in absolute value when the solution concentration increases. At 25°C, (8) provides b = -20 mV. This value is close to the one obtained by Pride and Morgan [1991], using a compilation of experimental data, who found b = -26 mV. The parameter c depends on temperature, on the total site density of the silica surface. and on the dissociation constant for

$$> SiOH^0 \quad \longleftrightarrow \quad > SiO^- + H^+$$
(9)



For volcanic rocks, having usually a complex mineralogical composition, the effect of pH and fluid conductivity on zeta potential cannot be explained analytically, using microscopic approach, as for quartz systems. The effect of these parameters may only be inferred empirically using experimental data. Unfortunately, few experimental works exist in the literature concerning the effect of these parameters on zeta potential. In particular, only two studies explored the effect of pH and fluid conductivity on zeta potential in rocks having mineralogical compositions similar to our samples: the Ishido and Mizutani [1981] work which examined the effect of pH in an andesitic sample and the Massenet and Van Ngoc [1985] work which studied the effect of pH and fluid concentration in samples of volcanic ashes (from Mount Etna). The mineralogical composition of these ashes is relatively close to the composition of our samples.

In order to calculate the values of the ζ potential of our samples, in the same physical conditions, we assumed that the variation of the zeta potential with water concentration and pH can be described by a relation similar to that of (8), as for quartz systems. We used the measurements of Massenet and Van Ngoc [1985] at $6 \le pH \le 7$ and fluid concentration $\le 10^{-3}$ mol L⁻¹, to deduce the values of b and d parameters. We used these values (b = 0.4 mV and d = 8 mV) to calculate zeta potential of our samples at $C_f = 2.1 \times 10^{-5}$ mol L⁻¹ and pH = 7. The values of zeta potential at $C_f = 2.1 \times 10^{-5}$ mol L⁻¹ and pH = 7, after pH and fluid concentration corrections (equation (8)), are given in Table 3. Figure 8 shows the zeta potential as a function of permeability before and after correction. We observe that after both corrections the dispersion of zeta potential is considerably reduced. The zeta potential ranges now from -4 to -19 mV. The zeta potential reflects differences in mineralogical composition of the samples. It is therefore related to the different eruption mechanisms, Figure 7. Computed parameter C' versus permeability, and we can observe that the lower values of zeta poten-



Figure 8. Zeta potential (a) deduced from equation (1) and (b) calculated from equation (8).

tial are obtained in the brecchias and scorias of Paléoand Néo-Pelée stages, while the higher values are observed in the 1929 eruption. For the P1 and 1902 eruptions, zeta potential is more important in samples from Peléan mechanism. These observations show that the zeta potential is very sensitive to the weak mineralogical composition variations.

5. Streaming Potential on Volcanoes

SP surveys on mountainous areas usually show a negative correlation between electrical potential and topographic elevation, implying that the higher the topographic elevation, the lower the potential. Such a correlation between SP and topographic elevation is the result of steady state fluid flow caused by spatial variations in the elevation of the water table [Ishido, 1989].

Indeed, the electrokinetic coupling coefficient is usually negative for rocks, and positive charges are induced to move along the fluid flow from the top to the bottom of the topographic slope. This SP anomaly reflects the piezometric surface, and the amplitude of the anomaly is related to the thickness of the nonsaturated zone [Aubert and Dana, 1994]. This topographic effect in a volcanic area can be -1.6 mV m^{-1} [Hashimoto and Tanaka, 1995], -1.9 mV m⁻¹ [Lénat, 1987], -3 mV m⁻¹ [Aubert and Lima, 1986], -5 mV m⁻¹ [Aubert and Dana, 1994], or larger, -10 mV m^{-1} [Ishido, 1989; Hashimoto and Tanaka, 1995]. In volcanic zones this topographic effect is modified by ascendant fluid flow, so that positive anomalies are usually observed, although it is not unusual to measure positive anomalies on extinct volcanoes too, induced by higher water-repellent levels. Large positive SP anomalies have been reported on La Fournaise volcano [Zlotnicki et al., 1994], on Mount Somma-Vesuvius and Vulcano Island [Di Maio et al., 1997], and on Unzen volcano [Hashimoto and Tanaka, 1995], and these anomalies are interpreted as the electric potentials generated by fluid flow in the hydrothermal ascendant circulations in the subsurface or greater depths. Recently, a huge anomaly of 4000 mV in self potential was reported on Misti volcano (south Peru) [Finizola et al., 1998]. Since streaming potential is proportional to the pressure gradient (Figure 4), any source of fluid pressure gradient change will be an important factor for the electrokinetic process. Temporal monitoring of SP on volcanoes could reflect climatic variations [Morat et al., 1992], variations in hydrothermal circulations resulting from crack closure or opening, fluid pressure increases induced by stress changes, or thermal fluctuations [Malengreau et al., 1994].

Electrical resistivity of water sources and rivers on Mount Pelée has been measured to be $10-120 \Omega$ m, with pH = 6.4. Sources of hot water (up to 73°C) showed also pH = 6.4. Moreover, resistivity of aquifers, measured via boreholes, was 20-72 Ω m [*Barat*, 1984]. Considering an average water electrical resistivity of 50 Ω m, one could expect a coupling coefficient of about -25 to -406 mV MPa⁻¹ in this volcano.

6. Conclusion

The measured coupling coefficient on 11 samples from Mount Pelée volcano has been found to vary with permeability. This variation has been explained by the effect of the effective conductivity, of the pH, and of the fluid conductivity. The remaining variation of the zeta potential is due to the differences in mineralogy of the samples related to the eruption mechanisms.

This study demonstrates that it is important to perform electrokinetic measurements by controlling pH and ionic strength (or fluid conductivity) and by measuring the effective conductivity. A good understanding of the electrokinetic phenomenon on rock samples must involve joint measurements on intact samples and on crushed samples, in order to know the exact value of zeta potential in andesitic-water systems and its dependence on salinity and pH. The electrokinetic process in unsatured medium and the effect of temperature have been theoretically quantified recently [*Revil* et al., 1999b]. The effects of vapor, and other gases, specially CO_2 , on electrokinetics in volcanic samples, must be quantified in order to better understand the SP anomalies observed on volcanoes. Indeed, very recent measurements show that injected gases in soil, with various effects on pH, induced different SP responses [Martinelli, 1998].

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