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Article

Estimating the Impact of Inflow on the Chemistry of Two Different Caldera Type Lakes Located on the Bali Island (Indonesia)

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Abstract: This paper is a preliminary attempt to assess the composition of pollutants in two different caldera lakes situated in the Indonesian Archipelago: Batur and Bratan. Both lakes are characterized by largely different physico-chemical regimes; Batur Lake is located in an area that is currently volcanically active, unlike Bratan Lake. The latter is much smaller and shallower than Batur Lake. The concentration of pollutants in the Indonesian equatorial lakes is largely unknown, and the impact of both biological and geothermal processes on their distribution requires attention. This study shows that the concentrations of cations (Na⁺, K⁺, Mg²⁺, Ca²⁺), anions (SO₄²⁻, F⁻, Cl⁻, Br⁻) and trace elements (Li, B, Al, V, Cr, Mn, Co, Ni, Cu, Zn, As, Se, Rb, Sr, Mo, Cd, Sn, Cs, Ba, Pb, U,

Be, Ag, Sb, Tl, Bi) differ greatly between both lakes. Most chemical parameters determined in volcanically influenced Batur Lake were tens to hundreds times higher than in Bratan Lake, and in the case of trace metals, the ratios of the two lakes' concentrations reached several hundreds. This study also compared the composition and concentration levels of organic compounds in both lakes, such as fatty acids, halocarbons and esters. On the other hand, the content of organic chemicals in the lakes also results from biological activity by phytoplankton, zooplankton and bacteria.

Keywords: caldera type lake; water pollution; ions; trace elements; total organic carbon (TOC); fatty acids

1. Introduction

Volcanic lakes can range in salinity from very low to hyper-acidic-brines. Variations in lake-water chemistry are related to variations in the composition and flux of volcanic fluids and gases entering the lake. Furthermore, they are modified by the dilution and evaporation effects [1]. Caldera lakes are filled volcanic cavities of two different origins, closely corresponding to the prevailing rock types. The explosive eruptions of high-viscosity, silica-rich magma cause the volcanic cone to rupture, leaving a basin for a future lake. This is the usual origin of andesitic-dacitic, rhyolitic, and peralkaline calderas [1]. The basaltic lavas, on the other hand, are characterized by effusive eruptions [2], thus a caldera can only be formed in them following the collapse of an empty magma chamber [3,4].

On Bali, Indonesia, natural lakes are typically of volcanic origin, usually formed in calderas, like Batur and Bratan Lakes discussed here (Figure 1).

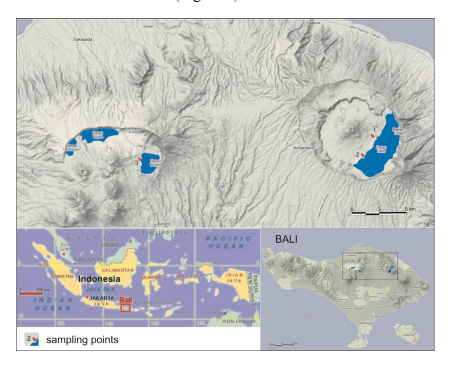


Figure 1. Location of the research area and water sampling points: 1—Batur Lake; 2—Batur Spring; 3—Bratan Lake (Source: www.istp.murdoch.edu.au).

Both calderas presented in Figure 1 host freshwater lakes but possess different physicochemical characteristics. Batur Lake (15.9 km², 88 m deep), is four times bigger than Bratan Lake, and is located 200 m lower. The Batur caldera is also volcanically active, having experienced 28 eruptions since 1800 AD, whilst the Bratan caldera has been dormant for at least 200 years (Figure 2). Only the volcanic activity of the nearby Agung caldera can influence the Bratan caldera, however this has been sporadic since 1840 AD.

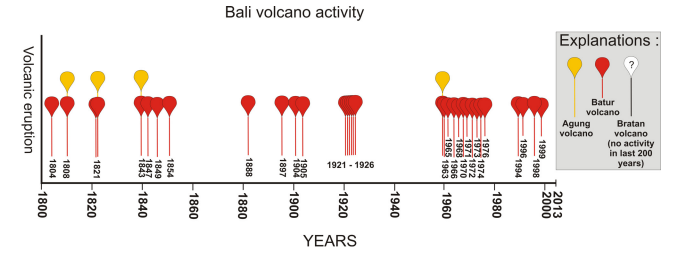


Figure 2. Volcanic activity of volcanoes situated near Batur and Bratan Lakes.

Batur Lake is located at the foot of the Batur volcano, which is 1717 m high and lies within two concentric, older calderas. These have diameters of 9 and 13 km, and are twenty and thirty thousand years old, respectively. The separate caldera of the Batur Lake, was formed in seven stages. The first pre-caldera reached a height of 3000 m above sea level and was composed either entirely of basalt or of basalt and andesite. The eruption of the second stage took place around 29,300 BC, introducing dacite and ignimbrite into the local substratum. During the third stage, the formation of the outer caldera occurred. The depression was deeper in the north-western part, causing outflow from an andesite magma reservoir and eruption of silica-rich lava in the fourth stage. The big eruption, alongside outflows of andesite magma through the slits and cracks, occurred during the fifth stage around 20,150 BC, and numerous domes were created. The formation of the inner caldera preceded the emergence of the lake, which took place during the sixth stage. The seventh stage included the formation of a stratovolcano of basalt and andesite in the central part of the caldera [5].

The Buyan-Bratan caldera, located in north-central Bali, is a volcanic complex older than the Batur caldera. The diameter of the structure is around 10 km. There are three lakes and several volcanic cones of the stratovolcanic type inside the inner part of the caldera. The cones are topped with craters which have erupted at least once. Geological interpretation of Buyan-Bratan caldera formation is impeded by its surface being covered with sediment from earlier eruptions of the nearby Batur volcano. One of the few existing pre-caldera Buyan-Bratan sediment exposures is located in the vicinity of the Gitgit waterfall. The sediments are represented by congealed lava with tufas and breccias. In the other part of the caldera there are various rocks represented by basalt and andesite in the form of volcanic sand and gravel breccias. The youngest pumice rocks were created before 23,000 BC. For over 200 years, volcanic eruptions in the caldera have not been recorded [6].

The lakes formed in the two calderas are likely to have contrasting characteristics, and we based our sampling methodology for both lakes upon this assumption. Due to the limited time available for conducting this fieldwork, the number of samples was limited to twelve from the caldera lakes and the adjacent spring. However, additional insight into the chemistry of these lakes can be gained from archived data obtained in 1992 and 1993 [4,7]. The sampling sites were selected following a precise morphological analysis of the area, and past chemical and hydrological records.

This paper is a thorough comparison of the chemical composition of these different caldera lakes, which represent different geomorphological characteristics. Since the chemistry of Indonesian equatorial lakes is largely unknown, this study will offer insights into the influence of geothermal processes on lake water chemistry. Furthermore, we also conducted an analysis of organic compounds to distinguish those originating from local biological activity from the typical organic pollutants.

This paper contains the results of the main ionic elements analysis (cations: Na⁺, K⁺, Mg²⁺, Ca²⁺, and anions: SO₄²⁻, F⁻, Cl⁻, Br⁻), as well as trace elements (Li, B, Al, V, Cr, Mn, Co, Ni, Cu, Zn, As, Se, Rb, Sr, Mo, Cd, Sn, Cs, Ba, Pb, U, Be, Ag, Sb, Tl and Bi). These will be used to confirm the difference in the water source of these volcanic lakes.

2. Experimental

pyroxene, magnetite and olivine.

2.1. Field Site

The research area, Bali (8°25' S, 115°11' E, 563.3 km²), is located in the Indonesian Archipelago within the Lesser Sunda island group. Two large calderas (Mount Batur 1717 m a.s.l.; Mount Sangiyang 2093 m a.s.l.) are one of the most prominent landforms in the chain of volcanic mountain ranges on Bali. Batur Lake (17.18 km²) and Batur Spring (which are parts of the freshwater lake system of the Mt. Batur caldera) and also Bratan Lake (one of the unconnected lakes situated beside the Mt. Sangiyang caldera), were selected for the study pertaining to the concentration levels of pollutants in this volcanic area located in a tropical climate (Figure 1). Table 1 presents detailed data related to differences between calderas, which need to undergo further discussion.

Table 1. Geochemical and petrological data of rocks and lakes characteristics [3,7,8].

Bratan Caldera	Batur Caldera		
Morphological	l characteristics		
1231 m above sea level, area 3.8 km ² ,	1031 m above sea level, area 15.9 km ² ,		
maximum depth 22 m	maximum depth 88 m		
Volcani	c activity		
Earlier than around 500 ka. BP	Around 500 ka. BP		
No information about volcanic eruptions	28 volcanic eruptions (from 1804 to 2000)		
Geochemistry and	l petrology of rocks		
 The caldera is composed of basalt and andesite containing 45%-63% silica (SiO₂); The outer caldera zone is composed with dacite and rhyolite containing 65%-72% silica; Accumulation of minerals such as plagioclase, 	 The caldera is composed of basalt and andesite; The erupting magma is of two types, either basaltic or rhyolitic; The magmatic processes are rapid and likely 		

connected to a shallow supply system.

Table 1. Cont.

Bratan Caldera	Batur Caldera
Lake cha	racteristics
• The shallowest lake on Bali;	
strongly influenced by recreational activities,	 The largest and deepest lake on Bali;
motor boating and other outdoor activities,	• Small-scale agriculture around the lake, and an
hotels and restaurants;	increasing number of guest houses are being
 Some small-scale agriculture is maintained 	built; the lake is used for fisheries;
around the lake	 Thermo- and chemostratified,
 Thermo- and chemostratified. 	• Total biomass: 2.4 mg·L ⁻¹ ;
• Total biomass: 4.4 mg·L ⁻¹ ;	• Transparency ranging 3.0–3.2 m.
• Transparency is 1.8 m.	

2.2. Sampling

All water samples were collected between 23 and 28 January 2013, in 250 mL plastic bottles for the determination of inorganic compounds and 500 mL dark glass bottles for organic compound analyses, tightly closed and transported to the laboratory in Gdańsk University of Technology, Poland. Upon arrival at the laboratory, the samples for organic compound determination were frozen.

2.3. Analytical Methods

A summary of the analytical techniques, used here to determine the inorganic ions, trace elements and total organic carbon (TOC), is provided in Table 2. The analytical procedures were validated against certified reference materials: Reference Material No. 409 (BCR-409, Institute for Reference Materials and Measurements, Belgium), ANALYTICAL-8 (Inorganic ventures, Christiansburg, VA, USA) and Analytical Reference Material TM-DWS.2 (Environment Canada).

Table 2. Validation parameters and technical specifications of the analytical procedures.

Parameter	Measurement Range	LOD	LOQ	CV (%)	Measi	urement Instrumentation	
Cations a,*	0.03-500	0.01	0.030	0.9	DIONEX 3000 chromatograph	column: Ion Pac® CS14 (3 × 250 mm); suppressor: CSRS-300, 2 mm, mobile phase: 38 mM metasulfonic acid, flow rate: 0.36 mL·min ⁻¹ , detection: conductivity	
Anions a,*	0.06–500	0.055-0.09	0.027–0.17	0.6	(DIONEX, Sunnyvale, CA, USA)	column: Ion Pac®AS22 (2 × 250 mm); suppressor: ASRS-300, 2 mm, mobile phase: 4.5 mM CO ₃ ²⁻ , 1.4 mM HCO ³⁻ , flow rate: 0.38 mL·min ⁻¹ , detection: conductivity	
TOC a,*	0.5-500	0.17	0.50	5.0	TOC analyzer (TO	OC-VCSH/CSN, SHIMADZU, Japan)	
Metals **	0.002-1000	0.0007	0.002	0.5–1.5	Elan DRC, PerkinElmer, USA gas fed to the atomizer→Ar: 0.98 L·min ⁻¹ , plasma gas→Ar: 15 L·min ⁻¹		

Notes: * $(mg \cdot L^{-1})$; ** $(\mu g \cdot L^{-1})$; a the measuring range can be extended towards higher concentration ranges LOD—Limit of detection; LOQ—Limit of quantification; CV—Coefficient of variance.

Comprehensive Two-Dimensional Gas Chromatography coupled with Time-of-Flight Mass Spectrometry (GCxGC-TOF/MS) was used to identify a broad spectrum of organic compounds. The analytical procedure of organic compounds' identification consisted of the following steps. After water samples from both lakes were thawed, 500 cm³ of water was taken for examination and mixed with 10 cm³ of dichloromethane (DCM). The extraction was carried out for 2 h, after which the DCM phase was separated from the water phase, to be evaporated to a volume of 0.3 cm³. Its subsample of 2 µL volume was injected into the GCxGC-TOF/MS system (LECO, St. Joseph, MO, USA), to enable an analysis with the operating parameters listed in Table 3.

Table 3. The operating parameters of the Comprehensive Two-Dimensional Gas Chromatography coupled with Time-of-Flight Mass Spectrometry (GCxGC-TOF/MS) system.

Inlet	Carrier Gas: Helium Inlet Mode: Splitless				
	GC1: 30 m ZB-5ms, film thickness 0.25 µm, inner diameter 0.25 mm.				
Chromatography columns	GC2: 1.5 m BPX50, film thickness 0.10 µm, inner diameter 0.1 mm.				
	Postcolumn: 0.21 m BPX50, film	thickness 0.10, inner diameter 0.1 mm			
	GC1:	GC2:			
	Target Temp (°C): 40 (Initial)	Target Temp (°C): 70 (Initial)			
Oven temperature	Duration (min): 1.0	Duration (min): 1.0			
Oven temperature	Rate (°C/min): 7.0	Rate (°C/min): 7.0			
	Target Temp (°C): 230	Target Temp (°C): 260			
	Duration (min): 10	Duration (min): 10			
	Modulation Period (s): 5.0				
Modulation parameters	Hot Pulse Time: 1.0				
	Cool Time Between Stages: 1.5				
	Start Mass (u): 33				
Mass speatromater settings	End Mass (u): 400				
wass spectrometer settings	Mass spectrometer settings Acquisition Rate (spectra/second): 200				
	Ion Source Temperature (°C): 250				
Library	NIST 2011, contains 243,893 spectra of 212,961 compounds.				

3. Results and Discussion

The specific physical processes influencing the chemical composition of Indonesian equatorial lakes are largely underexplored. The general setting suggests a stability of wind directions for prolonged periods and the likely important role of biological and geothermal processes [7]. The lake density structure depends on the gradients of total dissolved solids since temperature gradients are relatively small. Biologically induced stratification preserved by the salinity gradient seems to be of great importance, as well as deep thermal gradients which may be adiabatic and stable. Geothermal heating is likely to exceed the adiabatic gradient. Hot springs are likely to flow into the lakes, and in the case of Batur Lake, there is a thermal Batur spring. Seepage, rainfall and evaporation are their main sources of water supply. Changes in the mixing depth caused by the weather and depth of seasonal mixing may influence the nutrient flux into the epilimnion zone. Incomplete vertical mixing of lakes is highly dependent on the relative depth (or area) and how exposed they are to wind [7].

Tables 4 and 5 present the results of the research.

Table 4. Average concentrations of main elements analyzed in water samples from Batur Lake, Thermal Batur Spring and Bratan Lake [4,7].

Determined Parameter (mg·L ⁻¹)		Batur I	Lake	Batur Spring	Bra	atan Lake	Major Reservoirs in Indonesia
Year	2013	1992	1993	2013	2013	1993	1989–1995
pН	8.54	8.5	8.8	7.37	7.00	-	6.8-8.8
EC 1 (μ S/cm)	1,730	-	1.750-1.800	1,609	27.9	22–27	22-1,811
Na^+	287	355	350	248	3.40	1.3	1.3-350
K^+	55.7	31	22	55.3	0.388	0.45	0.45 - 22
Mg^{2+}	26.9	68	0.62	23.8	1.87	0.85	0.87 - 1,280
Ca^{2+}	18.2	35	35	29.6	0.082	1.9	1.9-32
$\mathrm{SO_4}^{2-}$	394	491	650-670	287	1.85	0.35 - 0.50	0.35-49,350
F^-	2.71	0.86	-	2.73	0.504	-	-
Cl ⁻	165	237	225	144	1.15	1.5	1.0-26,700
Br^-	0.723	-	-	0.881	0.118	-	-
Total N	0.22	=	0.256-0.970	3.39	0.030	0.310-1.310	0.116-1.310
Total P	2.03	-	0.028	2.08	0.360	0.002	0-0.080

Note: ¹ EC—Electrical conductivity.

Table 5. Average concentrations of trace elements analyzed in water samples from Batur Lake, the thermal Batur Spring, and Bratan Lake and rocks [7,9,10].

Determined Parameter (μg·L ⁻¹)	Batur Lake 2013	Batur Spring 2013	Batur 1992	Batur 1993	Batur Rocks (ppm)	Bratan Lake 2013	Major Reservoirs in Indonesia
Li	423	361	400			<lod< td=""><td></td></lod<>	
В	1440	2035	1490			53	
Al	2.4	1.1				8.0	
V	34.8	80.0			5.7-309	0.4	
Cr	3.8	6.2		$n.d.^1$	1.5-29	0.5	n.d. ¹
Mn	0.4	<lod< td=""><td></td><td></td><td></td><td>0.4</td><td>0-0.14</td></lod<>				0.4	0-0.14
Co	<lod< td=""><td><lod< td=""><td></td><td></td><td>2-45</td><td><lod< td=""><td></td></lod<></td></lod<></td></lod<>	<lod< td=""><td></td><td></td><td>2-45</td><td><lod< td=""><td></td></lod<></td></lod<>			2-45	<lod< td=""><td></td></lod<>	
Ni	<lod< td=""><td><lod< td=""><td></td><td>$n.d.^1$</td><td>2.0-43</td><td><lod< td=""><td>n.d.¹</td></lod<></td></lod<></td></lod<>	<lod< td=""><td></td><td>$n.d.^1$</td><td>2.0-43</td><td><lod< td=""><td>n.d.¹</td></lod<></td></lod<>		$n.d.^1$	2.0-43	<lod< td=""><td>n.d.¹</td></lod<>	n.d. ¹
Cu	2.2	2.6		$n.d.^1$	7.4-139	0.4	n.d. ¹
Zn	1.7	2.3		0-0.34		1.0	0-0.30
As	4.1	29.2				0.2	
Se	0.94	3.80				<lod< td=""><td></td></lod<>	
Rb	62.7	67.9			18-45	3.33	
Sr	96	140	1500		168-484	11	
Mo	3.15	5.76			0.6 - 2.2	<lod< td=""><td></td></lod<>	
Cd	<lod< td=""><td><lod< td=""><td></td><td>$n.d.^1$</td><td></td><td><lod< td=""><td>n.d.¹</td></lod<></td></lod<></td></lod<>	<lod< td=""><td></td><td>$n.d.^1$</td><td></td><td><lod< td=""><td>n.d.¹</td></lod<></td></lod<>		$n.d.^1$		<lod< td=""><td>n.d.¹</td></lod<>	n.d. ¹
Sn	<lod< td=""><td>0.50</td><td></td><td></td><td></td><td><lod< td=""><td></td></lod<></td></lod<>	0.50				<lod< td=""><td></td></lod<>	
Cs	3.13	4.29			0.3 - 2.5	0.04	
Ba	11.2	4.70			168-504	2.2	
Pb	<lod< td=""><td>< LOD</td><td></td><td>$n.d.^1$</td><td>3.3-12.9</td><td><lod< td=""><td>n.d.¹</td></lod<></td></lod<>	< LOD		$n.d.^1$	3.3-12.9	<lod< td=""><td>n.d.¹</td></lod<>	n.d. ¹
U	0.113	0.474			0.4 - 1.7	<lod< td=""><td></td></lod<>	
Be, Ag, Sb, Tl, Bi	<lod< td=""><td><lod< td=""><td></td><td></td><td></td><td><lod< td=""><td></td></lod<></td></lod<></td></lod<>	<lod< td=""><td></td><td></td><td></td><td><lod< td=""><td></td></lod<></td></lod<>				<lod< td=""><td></td></lod<>	

Note: n.d.1—not detected.

3.1. Main Elements

The results obtained with the use of analytical techniques (including GCxGC) indicate striking differences in the chemical composition of the water between the two lakes. The chemical parameters of the water samples obtained during the study are listed in Table 4, combined with data obtained from literature. The electrical conductivity (EC) in natural Indonesian lakes of varying origin (caldera, tectonic, tectonic/volcanic, floodplain, semi-natural, landslide, volcanic/crater, crater) was usually within the range of 80–300 μ S·cm⁻¹. Generally, the EC value decreases slightly with the increasing depth of a lake (Figure 3). However, this is not obvious for all caldera lakes. In Batur and Bratan Lakes, there were two extremes observed: 1730 and 27.9 μ S·cm⁻¹, respectively. Such values were earlier observed in 1993 for Batur Lake (1750–1800 μ S·cm⁻¹) and for Bratan Lake (22–27 μ S·cm⁻¹) [7]. Particularly low EC values observed in Bratan Lake may result from a short retention time of rainwater supplying this lake.

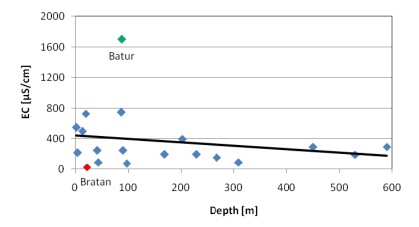


Figure 3. Batur and Bratan within the spectrum of electrical conductivities (EC) experienced in Indonesian lakes.

The concentration levels of the main ions and conductivity of Batur Spring water are very similar to Batur Lake water. The slope coefficient of a regression curve (between Batur Lake and Batur Spring) is close to one (0.926) for a very high correlation coefficient R² (0.997), the linear regression parameters of ions (Na⁺, K⁺, Mg²⁺, Ca²⁺, SO₄²⁻, F⁻, Cl⁻ and Br⁻) concentrations and electrical conductivity in the three studied sites (Table 6). A similar EC value (1609 µS·cm⁻¹) appears for the thermal Batur Spring. Batur Lake is both thermo- and chemostratified, although its studies were conducted only to the depth of 50 m. It is the most saline (165 mg·Cl⁻L⁻¹) of the studied lakes and has a high pH of 8.54 (pH = 8.8 in 1993 [7]), especially in the epilimnion zone. The research literature shows that pH of the epilimnion zone in the Indonesian lakes ranged from 6.8 to 8.8 [9]. The usually high pH values also affect carbon dioxide equilibria, probably due to a relatively high daytime algal production. The values of pH for Bratan Lake and Batur Spring differ significantly (7.00 and 7.37, respectively). Bratan Lake is also thermo- and chemostratified but its water is the most diluted of all the Indonesian lakes. Its salinity at the level of 1.15 mgCl⁻·L⁻¹ is 100 times lower than in Batur Lake and the thermal Batur Spring. In Indonesia, the concentration of chloride ions in lake water can fall within the range of 1.5 to 225 mg·L⁻¹ and largely depends on their origin, and thereby the depth and surface area of a lake [7].

Table 6. The linear regression parameters of Na⁺, K⁺, Mg²⁺, Ca²⁺, SO₄²⁻, F⁻, Cl⁻ and Br⁻ concentrations and electrical conductivity in the studied sites.

Compared Data	Regresion Curve	R^2
Batur Lake/Batur Spring	y = 0.926x - 9.02	0.997
Batur Lake/Bratan Lake	y = 0.016x - 0.591	0.965
Batur Spring/Bratan Lake	y = 0.017x - 0.464	0.980

Alkaline cations, especially calcium and magnesium, appeared mainly as a result of the ion exchange processes with the magma creating the calderas, as a mechanism of weathering [3]. Calcium concentrations in the Indonesian lakes were usually between 1.9 and 35 mg·L⁻¹. The lowest concentration determined during the research was around 20 times lower in Bratan Lake (0.082 mg·L⁻¹) and the highest concentration was in Batur Lake (18.2 mg·L⁻¹) and Batur Spring (around 29.6 mg·L⁻¹). Magnesium concentrations in the Indonesian lakes ranged from 4.0 to 10.0 mg·L⁻¹. For both caldera lakes, the values of Mg²⁺ concentrations differ by an order of magnitude. In comparison to 1993, there was a significant increase in the content of these ions in the water of Batur Lake from 0.62 mg·L⁻¹ to 26.9 in 2013, whereas for Bratan Lake this value only doubled [7]. The increase in Ca²⁺ and Mg²⁺ concentrations in lake water can be connected with soil acidification, resulting in a higher export of these elements into water.

In the Indonesian lakes, potassium usually ranged from 0.5 to 3.0 mg·L⁻¹ and generally there were no distinct vertical differences. However, during the research conducted on the samples from Batur Lake and Spring, K⁺ was determined and the concentration of K⁺ amounted to 55 mg·L⁻¹, which is 150 times higher than in Bratan Lake. During the research from 1993, the K⁺ concentration for Batur Lake was 22 mg·L⁻¹ and then increased by 150%. K⁺ is an essential component of acidic effusive rocks which form calderas (rhyolite), but its increase can also be caused by the growing use of natural and mineral potassium fertilizers in the fields surrounding the lake. Sodium levels ranged from 3.4 mg·L⁻¹ (Bratan) to about 300 mg·L⁻¹ (Batur), while for other lakes in Indonesia these values were about 1.5–18.0 mg·L⁻¹ [7]. The research carried out in 2013 did not show any significant deviation from the published data for the years 1992–1993, which suggests the main ions to be of generally stable composition in the waters of both lakes.

Sulfate concentrations of the natural lakes are expected to be high due to volcanic surroundings. Minor eruptions and hot springs are permanent sources of sulfate influx in the area that should be particularly significant in Batur Lake, which is subject to current volcanic activity. The contrastingly high concentrations in Batur (650–670 mg·L⁻¹) and low concentrations in Bratan (0.35–0.50 mg·L⁻¹) confirmed this assumed pattern. However, since 1989, a continual decrease of SO₄²⁻ concentration to the present level of 394 mg·L⁻¹ has been observed in Batur Lake, which may be an indication of decreasing volcanic activity in this period. In most natural Indonesian lakes, concentrations ranged from 1.0 to 3.0 mg·L⁻¹ [7,10].

Nutrient concentrations in Indonesian lakes are typically low, especially in natural lakes, however a relative increase in their concentration is observed in deeper lakes. The concentrations in reservoirs are, surprisingly, at a similar level, despite agricultural and populated areas contributing to their catchments, but this may be due to wetlands and rice paddies acting as efficient nitrogen traps.

Similarly to nitrogen, the total phosphorus (total P) concentrations were rather low. This is also typical of the area since Indonesian soils are deficient in phosphorus. The highest total nitrogen (total N) value was observed in 1993 in Bratan Lake (1.310 mg·L⁻¹), whereas the highest total phosphorus concentration was only 0.085 mg·L⁻¹ and was noted in another Indonesian lake, named Ranau [7]. Generally, total phosphorus values in Indonesian lakes did not exceed 0.1 mg·L⁻¹ and slightly declined with the increasing lake depth (Figure 4). However, the conducted research indicates a significant increase in the total P value both in the Batur Lake caldera and the Batur Spring, within the 20 year period since the last survey. This may be a result of the growing use of surfactants (containing phosphates) in households (intensified tourism development). The observed variations in total P concentrations are likely to be due to soil acidification (fertilization) and expanding phosphorus export into water, which intensifies eutrophication. This is an important change, since in 1993 eutrophication in Indonesian lakes was a very rare occurrence [7].

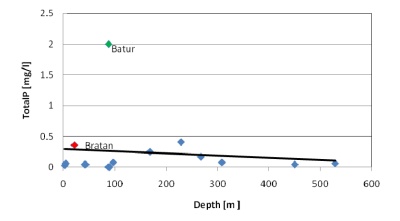


Figure 4. Relationship between the total phosphorus (total P) concentrations (highest values) and the depth in natural Indonesian lakes.

In 1993, the N:P-ratio for total nitrogen and total phosphorus in natural lakes was as expected, although they ranged from 1.9 to 485, and the particularly high ratios occurred due to low phosphorus concentrations. At present, high total P values make the N:P ratio 0.11 for Batur Lake and 0.083 for Bratan Lake, which is very low as compared to values from the 1993 survey, and indicates nitrogen limitation of their ecosystems. In this type of ecosystem, cyanobacterial blooms have a competitive advantage over other algal groups because of their nitrogen fixing abilities, and high P levels (>1 mg/L) may cause their intense blooms. This again follows from the important difference between the levels of phosphorus in these two lakes today, compared to an Indonesia-wide survey conducted in 1993, and is likely to be associated with current pollution. Therefore, nutrient regimes of Indonesian equatorial lakes should receive more attention, especially in respect to phosphorus.

3.2. Trace Elements

In 1993, heavy metals (cadmium, chromium, copper, nickel and lead) were not detected in Indonesian lakes. Only zinc concentrations were detected, and varied in natural lakes from undetectable to 0.34 mg·L⁻¹ in Batur Lake [7]. At present, the majority of metal ions, including heavy

metals, are detected and determined (Table 5). This is again a shift towards significantly higher pollutant levels than 20 years ago, which requires urgent attention from the local government.

The speciation of metals is modified by the environmental, physico-chemical factors such as pH, the presence of complexing ligands (humic and fulvic acids, chlorides, sulphates, carbonates or phosphates) and other metal ions. For the pH value of around 7 (for example in Bratan Lake) heavy metals occur in bioavailable and toxic forms. Humic and fulvic acids, chlorides, sulphates, carbonates or phosphates can also create stable complexes with metals. Such considerations on the influence of pH on the presence of trace elements have already been described for the Ijen caldera (Indonesia) [9].

Figure 5 shows the different characteristics of the analyzed trace elements, which can be related to their different sources and post-depositional paths, under conditions of various pH and organic compound concentration levels. The points circled in Figure 5 represent elements such as Cu, Zn, Mn, Cr, As, Se and U, which showed similar concentrations in both lakes. However, elements such as Rb and Sr, which occurred in high concentrations in Batur Lake, can be linked to volcanic activity (see Figure 2). From 1993 to 2013, the concentrations of these elements in Batur Lake have decreased slightly, which is consistent with the lack of fresh volcanic eruptions in the caldera during this period.

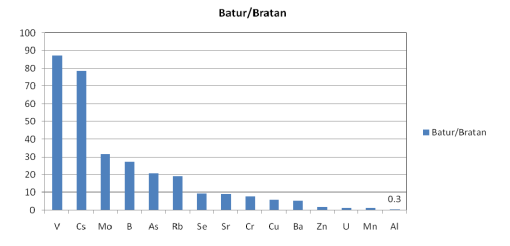


Figure 5. Relationship between the concentrations of chosen trace elements in both caldera lakes and the ratio of the given trace element concentration in Batur Lake to their concentration in Bratan Lake.

A decrease in the pH of lake water (for example from 8.8 to 8.54 at Batur Lake) influences the change in solubility and oxidation state of many metals. Under these mildly acidic conditions, the solubility of particularly Al, Cu, Zn, Ni and Pb has increased, while that of V decreased.

The bar chart in Figure 5 shows that the relationship between Al concentrations in both caldera lakes is significantly different from the relationships of other trace elements. The Al concentration for Bratan Lake is much higher ($8.0~\mu g\cdot L^{-1}$ for pH 7.00). The growing Al solubility can lead to the appearance of toxic ions, which may result in impoverishment of the local fauna and flora. Al can also precipitate in complexes with humic substances so that the water in a reservoir can become more transparent but harmful to animals, plants and human health.

Elements such as V, Cr, Co, Ni, Cu, Rb, Sr, Mo, Cs, Ba, Pb or U, which were detected and determined in the water of Batur Lake, are also present in volcanic gas plumes and rock obtained from the substrates (Table 2). The elements determined in the water of the lakes situated in the calderas also

contained minerals creating these calderas: Al (Plagioclase), Li (Pyroxene), V (Vanadium), Cr (Magnetite), and Ni (Olivine). Hydrothermal fluids, with element ratios that are different from those in host rocks, either precipitated secondary phases during their history, or were formed by an incongruent dissolution of volcanic rock during water–rock interaction. The main rock-derived elements in volcanic lake fluids from andesitic volcanoes are Na, Ca, K, Mg and Al, which are derived from the dissolution of volcanic glass, plagioclase, pyroxene and biotite or muscovite [10–12].

Table 7 presents the regression curve and correlation coefficients. The curve directional factor while comparing trace elements in Batur Lake and Batur Spring water is 1.37 for the high correlation coefficient R² (0.987). It is another indication that the basic inflow source for Batur Lake is Batur Spring water, with its high content of ionic and trace elements.

Table 7. Linear regression parameters of trace element (Li, B, Al, V, Cr, Mn, Co, Ni, Cu, Zn, As, Se, Rb, Sr, Mo, Cd, Sn, Cs, Ba, Pb, U) comparisons between field sites.

Compared Data	Regresion Curve	R^2
Batur Lake/Batur Spring	y = 0.926x - 9.02	0.997
Batur Lake/Bratan Lake	y = 0.016x - 0.591	0.965
Batur Spring/Bratan Lake	y = 0.017x - 0.464	0.980

3.3. TOC and Organic Compounds

Many publications hypothesize that the loss of dissolved organic matter from lake water is connected with their acidification, although the acidification itself may also be caused by the presence of humic compounds. Their concentration can be approximated using the total organic carbon (TOC) concentration levels [13–16].

A noticeable correlation has also been suggested between the TOC content and elevation above sea level [17]; the higher the altitude, the lower the TOC content in lakes. The TOC values for water from the two caldera lakes, differing in altitude by 200 m, is in stark contrast with the Batur Lake achieving a TOC concentration of 53.5 mg·L⁻¹ (Batur Spring reached 80.0 mg·L⁻¹ TOC), and Bratan Lake showing a relatively small TOC content of 4.09 mg·L⁻¹. The water of the higher situated, Bratan Lake, had a lower pH and TOC than the water of Batur Lake.

The total biomass amounts for Batur and Bratan Lake were 2.4 mg·L⁻¹ and 4.4 mg·L⁻¹, respectively, and these data correspond with the transparency of water (from 3.0 to 3.2 m for Batur and 1.8 m for Bratan). These small biomass values, and low nutrient levels, make the local ecosystem more vulnerable to small chemical changes [7]. Hence, the pollution with trace metals described here may be an important factor influencing the local biota. Interestingly, the total biomass in these lakes showed an opposite direction of changes to the TOC, perhaps implying that an important part of the TOC originates from human-induced pollution.

The analysis of DCM extracts from the caldera lake water was carried out with the use of a GCxGC-TOF/MS system. As a result of the separation of compounds contained in DCM extracts with a comprehensive two-dimensional gas chromatography, very detailed chromatograms were obtained (around 800–900 chromatographic peaks). These are presented in Figure 6, where parts A–D show intervals of retention time in the second column of the GcxGC-TOF/MS system.

These were cross-checked against a control sample (blank). A few of the peaks present in blanks also occurred in the samples, although their intensity was significantly higher than in the blanks. It may suggest that these compounds are also present in the sample. Aliphatic hydrocarbons, alcohols, ketones, aldehydes, organochlorides, organobromine and organoiodine compounds, and fatty acids were among the identified compounds. A considerable number of compounds (around 35%–40%) present such a complicated mass spectrum that their identification by means of comparing to spectrums from the NIST 2011 library does not allow us to define their structure with an appropriate probability. Table 8 shows a list of compounds identified in the samples which were not present in blanks.

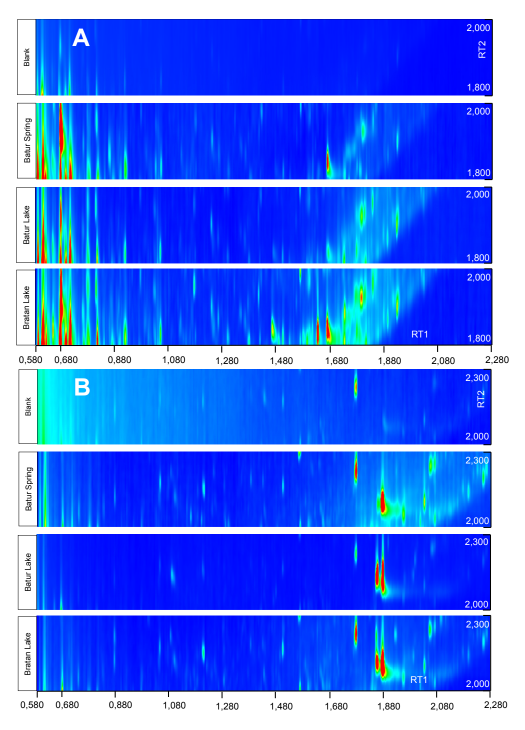


Figure 6. Cont.

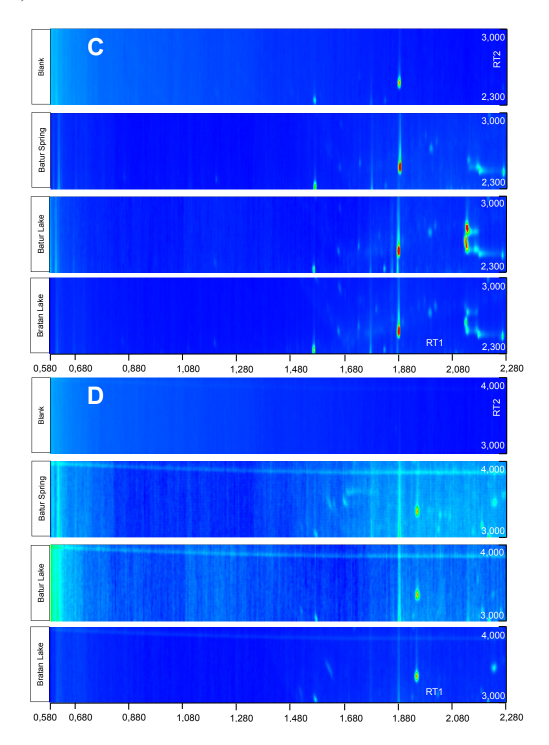


Figure 6. GCxGC-MS/TOF chromatograms of the dichloromethane (DCM) extracts, of the following samples from top to bottom: blank, Batur Spring, Batur Lake, Bratan Lake water. Sections A–D show intervals of retention time on the second column of the system: (**A**): 1800–2000 s; (**B**): 2000–2300 s; (**C**): 2300–3000 s; (**D**): 3000–4000 s. Several peaks visible in Figure 6 stem from the extract preparation protocol (the applied solvent—DCM, the employed glass, and the procedure of extract evaporation).

Table 8. A list of compounds identified in the samples which were not present in blanks.

Compounds Identified	Retention Time RT1; RT2	Blank	Batur Lake	Batur Spring	Bratan Lake
	RT: 1650–1800		Lake	Spring	Lake
Halocarbons		+	+++	+++	+++
	RT: 1800–2000				
2-(dodecyloxy)-ethanol	1630; 1830	-	+	+	+++
Tetradecanoic acid (Myristic acid)	1665; 1845	-	-	-	+++
trans-2-Undecen-1-ol	1725; 1845	-	-	-	-
	RT: 2000–2300				
5-butyldihydro-2(3H)-furanone	1115; 2140	-	+	-	-
cis-9-Hexadecenoic acid	1855; 2150	-	+++	+	+++
Hexadecanoic acid	1875; 2155	-	+++	+++	+++
Diethylene glycol monododecyl ether	1920; 2225	-	-	-	+
3-Heptadecanone	2070; 2260	-	-	+	+
	RT2: 2300-3000				
Dibutyl phthalate	1880; 2505	-	+++	+++	+++
2-methyl-Eicosane	1995; 2675	-	+	+	+
Oleic Acid	2130; 2550	-	+++	+	+
9,12,15-Octadecatrienoic acid	2135; 2710	-	+++	+	+
Dodecanamide	2210; 2990	-	-	+	-
Heptadecyl acetate	2265; 2460	-	+	+	+
<u>-</u>	RT2: 3000-4000				
Hexanedioic acid, mono-(2-ethylhexyl)-ester	1945; 3345	-	+	+	+

Notes: Compounds: not present in the sample(-); present in trace amounts(+); present at relativety high level(+++).

The broad profile of organic compounds found in both lakes was similar. The presence of fatty acids such as tetradecanoic acid (myristic acid, 14:0), hexadecanoic acid (palmitic acid, C16:0), octadecanoic (stearic acid, 18:0), 9Z-octadecenoic acid (oleic acid, 18:1), and 9,12,15-octadecatrienoic acid (linonenic acid, 18:3) is characteristic of the tested samples. The occurrence of fatty acids in the researched water is most probably connected with the presence of bacteria, including cyanobacteria, which are widely distributed in the free-living state and can be encountered in almost every habitat on the Earth. These bacteria are resistant to high salinity or acidity, prolonged droughts and high temperatures of hot springs. Cyanobacteria are able to produce a variety of fatty acids, but predominantly, it is the hexadecanoic acid, usually present at the level of 24%–40% of total fatty acids [18]. Hexadecanoic acid occurs in all tested samples collected from both Batur and Bratan Lakes and Batur Spring. Its concentration is relatively high and similar in all samples (Figure 7). There might be other sources of its origin, e.g., coconut palms.

Despite the general similarities, a range of organic compounds' concentrations distinguished the two lakes from one another. The deep waters of Batur Lake contain much higher oleic and linonenic acid concentrations than those observed in the waters of Bratan Lake. Myristic acid (14:0) was only identified in Bratan Lake and neither was observed in Batur Lake nor Batur Spring.

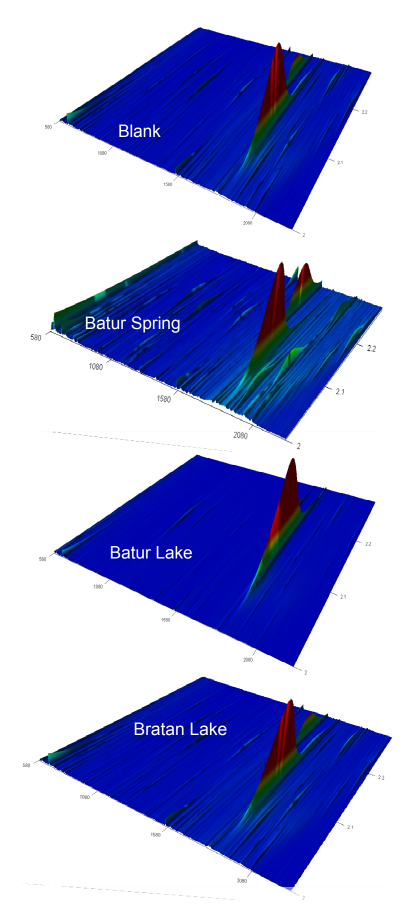


Figure 7. 3D chromatograms of blank and DCM extracts of Batur Spring, Batur, and Bratan Lake water in the retention time range for the second column from 2000 to 2300.

The differences in the fatty acid content and concentrations in both lakes may have resulted from a difference in the cyanobacteria species composition, a different content of dissolved CO₂ and nitrogen in the water of both lakes (also in the inflowing Batur Spring water), and the significant difference in their water ionic and trace metal composition. The more alkaline Batur Lake water, also deficient in CO₂, probably promotes the growth of cyanobacteria whose cell membranes include oleic and linonenic acids. At the same time, the poorly mineralized water of the Bratan Lake, with a neutral pH and similar N content, represents a different fatty acid profile, likely promoting the oleic and linonenic acids' biodegradation [18–21].

4. Conclusions

The results of our study on the abundance of major, minor, trace elements and organic compounds in the water from two different caldera lakes indicate that their chemistry is the result of important interactions among water, volcanic gases and the local substratum. In terms of pollution, the most important feature was the detection of multiple trace elements. The organic compounds were connected to local biota, and some of them could be used as distinguishing factors between the two lakes. The conducted research shows that water from both lakes significantly differs in cation (Na⁺, K⁺, Mg²⁺, Ca²⁺) and anion (SO₄²⁻, F⁻, Cl⁻, Br⁻) concentrations. The mentioned parameters are around 100 times higher in Batur Lake than Bratan Lake. Similar differences are also noted for conductivity and pH. The concentration levels of the mentioned ions and conductivity of Batur Spring water were very similar to the Batur Lake water. Their almost identical ionic composition of water strengthens the statement that this spring provides the main inflow supply for Batur Lake; however, a common groundwater source could not be excluded.

The concentration levels of trace elements in Batur Lake are found to be several to several hundred times higher than in Bratan Lake. However, the concentration levels in the water of Batur Lake and Batur Spring were very similar. On the other hand, the concentration levels of ions present in Bratan Lake are comparable to concentrations determined in the precipitate observed in the area of interest, as it is the main factor of loading water into this lake [22–24].

The presence of organic compounds in lakes results mainly from biological activity of microorganisms inhabiting them such as bacteria, plankton and zooplankton. The conducted research facilitates a comparison of the composition and level of fatty acid occurrence (the indicators of cyanobacteria presence) and the obtained results suggest diversity of microorganisms inhabiting both lakes.

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Author Contributions

Zaneta Polkowska designed the concept, selection of the study area, designed the experiment, analyzed and interpreted the data and wrote the draft of the manuscript, reviewed the final manuscript. Lidia Wolska provided his support and guidance throughout the field experiment, analyzed and interpreted the data and wrote the draft of the manuscript, reviewed the final manuscript. Leszek Łęczyński interpreted the results and prepared the manuscript, prepared figures. Marek Ruman selection of the study area, supervised data collection, helped with gathering field data, prepared figures, elaboration of geographic data and maps, reviewed the final manuscript. Sara Lehmann chemical and physical properties analysis. Katarzyna Kozak chemical and physical properties analysis, reviewed the final manuscript. Magdalena Matysik helped with gathering field data, supervised data collection, elaboration of geographic data and maps. Damian Absalon helped with gathering field data, supervised data collection, elaboration of geographic data and maps

Conflicts of Interest

The authors declare no conflict of interest.

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