

Amazon River dissolved load: temporal dynamics and annual budget from the Andes to the ocean

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Abstract The aim of the present study is to estimate the export fluxes of major dissolved species at the scale of the Amazon basin, to identify the main parameters controlling their spatial distribution and to identify the role of discharge variability in the variability of the total dissolved solid (TDS) flux through the hydrological cycle. Data are compiled from the monthly hydrochemistry and daily discharge database of the “Programa Climatológico y Hidrológico de la Cuenca Amazonica de Bolivia” (PHICAB) and the HYBAM observatories from 34 stations distributed over the Amazon basin (for the 1983–1992 and 2000–2012 periods, respectively).

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This paper consists of a first global observation of the fluxes and temporal dynamics of each geomorphological domain of the Amazon basin. Based on mean interannual monthly flux calculations, we estimated that the Amazon basin delivered approximately 272×10^6 t year⁻¹ (263–278) of TDS during the 2003–2012 period, which represents approximately 7 % of the continental inputs to the oceans. This flux is mainly made up by HCO₃, Ca and SiO₂, reflecting the preferential contributions of carbonate and silicate chemical weathering to the Amazon River Basin. The main tributaries contributing to the TDS flux are the Marañón and Ucayali Rivers (approximately 50 % of the TDS production over 14 % of the Amazon basin area) due to the weathering of carbonates and evaporites drained by their Andean tributaries. An Andes–sedimentary area–shield TDS flux (and specific flux) gradient is observed throughout the basin and is first explained by the TDS concentration contrast between these domains, rather than variability in runoff. This observation highlights that, under tropical context, the weathering flux repartition is primarily controlled by the geomorphological/geological setting and confirms that sedimentary areas are currently active in terms of the production of dissolved load. The log relationships of concentration vs discharge have been characterized over all the studied stations and for all elements. The analysis of the slope of the relationship within the selected contexts reveals that the variability in TDS flux is mainly controlled by the discharge variability throughout the hydrological year. At the outlet of the basin, a clockwise hysteresis is observed for TDS concentration and is mainly controlled by Ca and HCO₃ hysteresis, highlighting the need for a sampling strategy with a monthly frequency to accurately determine the TDS fluxes of the basin. The evaporite dissolution flux tends to be constant, whereas dissolved load fluxes released from other sources (silicate weathering, carbonate weathering, biological and/or atmospheric inputs) are mainly driven by variability in discharge.

These results suggest that past and further climate variability had or will have a direct impact on the variability of dissolved fluxes in the Amazon. Further studies need to be performed to better understand the processes controlling the dynamics of weathering fluxes and their applicability to present-day concentration–discharge relationships at longer timescales.

Keywords Amazon basin · Andes · Sedimentary areas · Large rivers · Water chemistry · Dissolved solid flux · Weathering · Hydrological variability

Introduction

Solutes in large rivers are the main tracers of the element cycles of the Earth's critical zone at the continental scale. The estimation of their annual fluxes and the description of their concentration variability in response to the hydrology allow the determination of the weathering budget of contrasted environments and their responses to climate forcing. Because of its geological and climate heterogeneity, the Amazon basin can be considered as a “natural laboratory” allowing, for example, for the evaluation of the role of these environmental factors in matter exportation budgets. Moreover, the riverine solute exports are also an important source of nutrients for the oceans. At the global scale, the tropical basins are the main source of dissolved matter for the oceans (Gaillardet et al. 1999; Meybeck 2003). Among these tropical basins, the Amazon River is a major source of dissolved solids (e.g. Milliman and Farnsworth 2011).

Historically, the waters of the Amazon streams were classified based on their colour, which is related to their hydrochemical characteristics (transparency, pH, conductivity, organic matter content and suspended matter) and to their ecological properties (e.g., Sioli 1964; Junk and Piedade 1997; Rios-Villamizar et al. 2014). These characteristics are in turn linked to the geological, geomorphological and biological properties of each sub-basin. The chemical composition of the main tributaries of the Amazon has allowed for the discrimination of the lithological sources of the dissolved load (Stallard and Edmond 1983, 1987; Edmond et al. 1996; Gaillardet et al. 1997; Mortatti and Probst 2003; Moquet et al. 2011). From these studies, it appears that the source of the dissolved load is mainly associated with silicate and carbonate weathering processes in the Andes. Nevertheless, Moquet et al. (2011) and Bouchez et al. (2012, 2014) suggested that the lowland sedimentary areas contribute to dissolved load production and that the CO₂ consumption associated with the weathering of floodplain sediments is significant for the Amazon basin budget. From these studies, it also appears that the Andes silicate weathering flux is nearly proportional to the water flux, at least at the annual timescale (Moquet et al. 2014a).

However, the methods used to discriminate the sources of dissolved load are associated with significant uncertainties that might obscure the variability of the dissolved load along the hydrological cycle. These uncertainties are mainly associated with the definitions of the silicate end member compositions (Gaillardet et al. 1997; Bouchez and Gaillardet 2014; Moon et al. 2014) and with the identification of the SO₄²⁻ sources (Calmels et al. 2007; Beaulieu et al. 2011), especially in the case of the Andean basins (Moquet et al. 2011). In the present study, we explore the major element concentrations and fluxes without applying any correction/discrimination, and we focus on (1) the relationships between solute production and water flux at a timescale shorter than 1 year and on (2) the identification of the geographical areas producing each major solute.

Although the total suspended solids (TSS) dynamics and budget in the Amazon have largely been documented in the literature, both at the scale of the whole Amazon basin (e.g. Meade et al. 1985; Dunne et al. 1998; Filizola and Guyot 2009; Martinez et al. 2009; Filizola et al. 2011) and of the Andes (Guyot et al. 1996; Armijos et al. 2013a, b; Pepin et al. 2013; Santini et al. 2014), the total dissolved solid (TDS) variability in response to the hydrological cycle has been less examined. Some local studies have been performed in the shields (Markewitz et al. 2001), in the central plain area (Devol et al. 1995), at the outlet of the Amazon (Oltman et al. 1964; Oltman 1967; Gibbs 1967a, b) and on Andean rivers (Roche and Fernandez Jauregui 1988; Guyot et al. 1993; Townsend-Small et al. 2008; Aufdenkampe et al. 2007; Wilcke et al. 2001, 2003; Boy et al. 2008; Armijos et al. 2013b; Torres et al. 2015). Some studies provide a general overview of the temporal variability of TDS (e.g. Mortatti and Probst 2003; Tardy et al. 2005; Bustillo et al. 2010, 2011; Sanchez et al. 2015), but these results are restricted to a limited number of hydrological stations, and the Andean rivers are not specifically considered. To date, no study has presented an integrated view of the TDS variation as a function of discharge from the Andean basins to the outlet.

The present paper is a complementary study of Moquet et al. (2011) and Sanchez et al. (2015) and includes the database previously used by these authors. These studies estimate the weathering budget of the Andean basins (Moquet et al. 2011) and report estimates of dissolved load exports at the scale of the main tributaries (Sanchez et al. 2015). In the present study, we expanded our analysis to all the available contents of the geodynamical, hydrological and biogeochemical control of erosion/alteration and material transport in the Amazon, Orinoco and Congo basins—Environmental Research Observatory (HYBAM observatory) and PHICAB databases (>3,000 samples analysed for major elements over 34 gauging stations) at the outlet of the main geomorphological domains of the Amazon basin. We exploited this database following three main objectives. The first objective is to discuss the spatial variability of major element concentrations as a function of lithological, geomorphological and climate parameters. The

second objective is to show the role of intra-annual hydro-climatic variability in the variability of TDS concentrations. The third objective is to determine the dissolved load fluxes exported by the Amazon basin and to identify their main geographical origin at the basin scale.

Study area

The Amazon basin covers 5.9 million km² (Callède et al. 2010). It is located between 5° 16' N and 20° 28' S, between 79° 36' W and 50° 15' W and across five main countries: Brazil (63 % of the surface area), Peru (16 %), Bolivia (12 %), Colombia (6 %) and Ecuador (2 %). Negligible portions of the basin are also included in Venezuela and Guyana. With an annual discharge of 6.5–6.6×10¹² m³ year⁻¹ (Dai and Trenberth 2002; Callède et al. 2010), the Amazon River contributes 16–18 % of the continental discharge to the oceans according to the global estimates of Dai and Trenberth (2002) and GRDC (2014). In the present work, the study area corresponds to the Amazon basin at the Óbidos station (Amazon River), Itaituba station (Tapajós River) and Altamira station (Xingu River) and covers 5.6 million km², representing 94 % of the entire Amazon basin (Fig. 1).

The Amazon basin can be separated into three geomorphological domains (Fig. 1 and Table 1):

1. The Andes (25 % of the basin) is the active orogenic zone and the major source of dissolved matter (due to the abundance of evaporite and carbonate outcrops; Gibbs 1967b; Stallard and Edmond 1987) and of suspended solids (e.g. Guyot et al. 1996; Armijos et al. 2013a, b). In the present study, this domain includes the Andean forelands (approximately half of this area), corresponding to Cenozoic back-arc basins (Roddaz et al. 2005), which make up the transition zone between the Andes and the central plain. This domain can behave as a sediment trap or as an erosional surface, according to the regional structural dynamics (Guyot 1993; Laraque et al. 2009; Baby et al. 2009; Armijos et al. 2013a, b; Santini et al. 2014).
2. The central plain domain (also named ‘lowland’; 24 % of the basin) forms a sedimentary basin for inputs coming from the Andes and, in much lower proportions, from the shields. It corresponds to the zone occupied by the main outflow channel of the Amazon toward the ocean (Filizola et al. 2011). This area is punctuated by floodplain lakes (*várzeas*) along the main channel of the Solimões and Amazon Rivers, which respond to the seasonal regime

Fig. 1 Topographic map of the Amazon basin and locations of the monitored gauging stations of the PHICAB program and HYBAM observatory (Gauging stations are detailed in Table 2). (*M.d.D.* Madre de Dios)

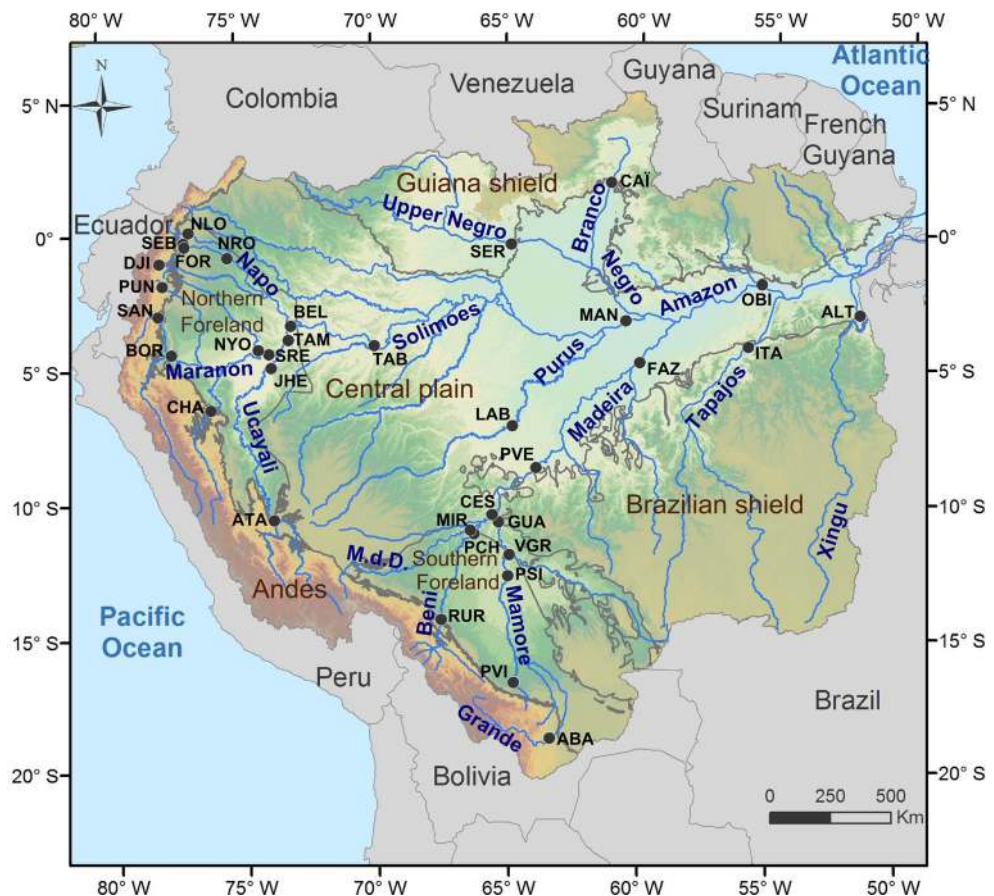


Table 1 Amazon basin and geomorphological domains (% of the total Amazon basin area)

	Geomorphological setting							Total
	Andean domain			Central plain	Shields			
	Andes (>400 m)	Andean forelands	Total		Brazilian shield	Guyana shield	Total	
Area (10 ⁶ km ²)	683	779	1,462	1,452	853	2,186	3,039	5,953
% total Amazon basin	12 %	13 %	25 %	24 %	14 %	37 %	51 %	100 %

See Fig. 1 for the definition of domain boundaries

of inundation. These areas can play a significant role in the geochemical cycle of the Amazon sedimentary plain (Viers et al. 2005; Cullmann et al. 2006).

3. The Guiana and Brazilian shields (51 % of the basin) lie to the north and south of the Amazon corridor. They consist of two extensively eroded Precambrian terrains and have little influence on the inorganic dissolved and solid fluxes carried by the Amazon (Furch et al. 1982; Konhauser et al. 1994).

The hydrological network of the Amazon basin is organized according to five main tributaries: the Negro, Solimões, Madeira, Tapajós and Xingu Rivers. The following water budget and areas are compared with the corresponding number after the confluence of the Amazon, Tapajós and Xingu Rivers. It is based on the 2003–2012 discharge at the HYBAM observatory database at the outlet stations of these rivers reported in Table 2 and by Getirana et al. (2010) for the Negro River budget (1997–2006 period). The Negro River, near Manaus City (~17 % of the Amazon discharge and 13 % of its area; Getirana et al. 2010), in the north of the basin, drains a large part of the Guyana shield. The main tributaries are the Branco River to the east and the Upper Negro River to the west. The Solimões River (51 % of the Amazon annual discharge and 39 % of its area) drains the Northern and Central Andean sub-basins and the main part of the central plain. The main Andean Solimões tributaries are, from north to south, the Napo, Marañón and Ucayali Rivers. The Napo basin includes most of the active volcanoes of the basin. The Marañón and the Ucayali drain diverse lithologies and are especially marked by the occurrence of karstified carbonates and evaporite domes (Stallard and Edmond 1983). After the confluence of the Napo, Marañón and Ucayali, the main left bank tributaries of the Solimões are the Iça and Japura Rivers that drain the northern central plain and the small Colombian Andean basins. The main right bank tributaries are the Javari, Jutá, Juruá and Purus Rivers, which drain the centre part of the central plain. The Madeira River (13 % of the Amazon discharge and 23 % of its area) drains the southern Andes and the western part of the Brazilian shield. The main Andean tributaries of the Madeira River are, from north to south, the Madre de Dios, the Beni and

the Mamoré. Mamoré basin includes the semi-arid basin of the Grande River. The Tapajós and Xingu Rivers (6 and 4 % of the Amazon discharge, respectively, and 8 and 8 % of its area) drain the remaining area of the Brazilian shield in the south-southeastern part of the basin. These five main Amazon tributaries (Negro, Solimões, Madeira, Tapajós and Xingu) connect with the Amazon River in the central Amazon plain between the cities of Manaus and Belem (Brazil) from ~1,200 to 150 km upstream from the estuary.

The climate of the Amazon basin is monsoonal and is controlled by the Southern American Monsoon System (SAMS). It depends on the seasonal variation of continent-ocean contrast in temperature and the seasonal migration of the intertropical convergence zone (ITCZ) southward (northward) during austral summer (winter) (Marengo 2004; Marengo et al. 2012). The ITCZ causes deep atmospheric convection with high rainfall and contributes a constant supply of moisture from the Atlantic Ocean to the Amazon Basin (Nogués Paegle et al. 2002; Vera et al. 2006). The Andean mountain chain acts as an orographic barrier and affects the atmospheric circulation and rainfall patterns at the continental scale (Insel et al. 2009; Garreaud et al. 2009). This orographic effect also induces a high heterogeneity of the rainfall distribution and intensity over the Andes and can favour either extremely humid (hotspots) or extremely dry (semi-arid) conditions in this region (Vera et al. 2006; Garreaud et al. 2009; Espinoza et al. 2009a).

At the scale of the basin, a north–south gradient is observed in the precipitation amount and seasonality (Espinoza et al. 2009a, b). Northern and northwestern Amazonia are exposed to high rainfall associated with a low seasonality, whereas southwestern and southern Amazonia experience lower rainfall associated with high seasonality and a dry (humid) season during austral winter (summer). This affects the hydrographs (Guimberteau et al. 2012) (Fig. 2). The maximum monthly discharges of the Tapajós, Madeira, Solimões and Negro Rivers occur in March, April, June and July, respectively. This time lag in maximum discharge reflects the south–north displacement of the ITCZ throughout the year (Fig. 2a) and is accompanied by a south–north gradient in the intensity of seasonality (the relative difference between the higher and lower monthly discharges) and in the specific discharge

Table 2 Geographical and hydrological characteristics of gauging stations

Hydrosystem	Sub-basin	Station name	River	Geomorphological setting	Observation network	Coordinates		Elevation (m)	Annual discharge ($10^3 \text{ m}^3 \text{ s}^{-1}$)	Area (10^2 km^2)	Water sampling period (number of sampling)
						Long. (°)	Lat. (°)				
Solimões	Napo	NLO	Aguarico	Andes	refHYBAM	-76.8083	0.044	300	0.5	5	2000–2008 (119)
		DJI	Jatunyacu	Andes	refHYBAM	-77.9186	-1.0833	640	0.3	3	2000–2008 (126)
		FOR	Napo	Andes	SO-HYBAM	-76.9784	-0.4737	260	1.3	12	2000–2012 (192)
		SEB	Coca	Andes	refHYBAM	-77.0068	-0.3429	270	0.4	5	2000–2008 (125)
		NRO	Napo	Andes + foreland	refHYBAM	-75.3964	-0.9169	180	2.2	27	2000–2008 (107)
		BEL	Napo	Andes+foreland	refHYBAM	-73.0761	-3.4783	90	6.5	100	2001–2011 (82)
		SAN	Santiago	Andes	refHYBAM	-78.0138	-3.0515	270	1.4	24	2001–2008 (110)
	Marañon	BOR	Upper Marañon	Andes	SO-HYBAM	-77.5482	-4.4704	150	5	114	2003–2012 (162)
		PUN	Pastaza	Andes	refHYBAM	-77.8258	-1.9141	660	0.7	13	2000–2005 (49)
		CHA	Huallaga	Andes	refHYBAM	-76.1193	-6.5704	180	2.9	69	2003–2010 (81)
		NYO	El Tigre	foreland	refHYBAM	-74.3823	-4.162	120	2	43	2003–2009 (55)
		SRE	Marañon	Andes + foreland	refHYBAM	-73.9081	-4.5162	100	17	357	2003–2011 (96)
		ATA	Ucayali	Andes	SO-HYBAM	-73.8179	-10.6782	210	6.1	191	2003–2011 (98)
		JHE	Ucayali	Andes + foreland	refHYBAM	-73.8261	-5.0273	110	12	353	2002–2011 (68)
Amazonas/Solimões	TAM	Amazonas	Andes + foreland	refHYBAM	-73.1617	-4.0031	90	30	722	2002–2011 (105)	
	TAB	Amazonas/Solimões	Andes + foreland	SO-HYBAM	-69.9603	-4.2418	70	35	880	2003–2011 (119)	
	LAB	Purus	Central plain	SO-HYBAM	-64.8	-7.2522	70	5.5	227	2004–2011 (76)	
	MAN	Solimões	Andes + foreland + central plain	SO-HYBAM	-60.5533	-3.3452	20	104	2,203	2003–2011 (86)	
	Madeira	VGR	Itenez	Brazilian shield	PHICAB	-64.93	-12.02	140	1.9	354	1982–1990 (128)
		ABA	Grande	Andes	PHICAB	-63.4095	-18.9091	450	0.4	59	1982–1990 (70)
		PVI	Mamore	Andes	PHICAB	-64.7867	-16.8409	210	0.5	8	1982–1990 (92)
		PSI	Mamore	Andes + foreland	PHICAB	-64.99	-12.81	140	4.3	234	1982–1990 (121)
		GUA	Mamore	Andes + foreland + Brazilian shield	PHICAB	-65.3458	-10.8078	120	7.9	618	1982–1990 (70)
		RUR	Beni	Andes	SO-HYBAM	-67.5343	-14.441	210	2.2	70	2003–2012 (103)
Beni	PCH	Beni	Andes + foreland	PHICAB	-66.26	-11.25	130	3.1	114	1982–1990 (59)	
	MIR	Madre de Dios	Andes + foreland	PHICAB	-66.4159	-11.1123	130	5.6	124	1982–1990 (88)	
	CES	Beni	Andes + foreland	PHICAB	-65.5812	-10.5347	110	9.8	276	1982–1990 (76)	
	PVE	Madeira	Andes + foreland + Brazilian shield	SO-HYBAM	-63.9212	-8.7877	60	18	954	2003–2012 (107)	
	FAZ	Madeira	Andes + foreland + Brazilian shield	SO-HYBAM	-60.0257	-4.8973	20	28	1,312	2003–2012 (113)	
Negro	SER	Upper Negro	Guyana shield	SO-HYBAM	-64.8263	-0.4974	30	19	290	2003–2011 (94)	
	CAI	Branco	Guyana shield	SO-HYBAM	-61.1227	1.8192	40	2.9	125	2003–2012 (108)	

Table 2 (continued)

Hydrosystem	Sub-basin	Station name	River	Geomorphological setting	Observation network	Coordinates		Elevation (m)	Annual discharge ($10^3 \text{ m}^3 \text{ s}^{-1}$)	Area (10^3 km^2)	Water sampling period (number of sampling)
						Long. (°)	Lat. (°)				
Amazon	Amazon	OBI	Amazon	Amazon basin	SO-HYBAM	-55.5111	-1.9471	10	178	4,669	2003–2012 (125)
Tapajós	Tapajós	ITA	Tapajós	Brazilian shield	SO-HYBAM	-55.9833	-4.2833	10	12	458	2004–2012 (90)
Xingu	Xingu	ALT	Xingu	Brazilian shield	discrete sampling	-52.0167	-3.0792	60	7.8	457	1997–1998 (2)

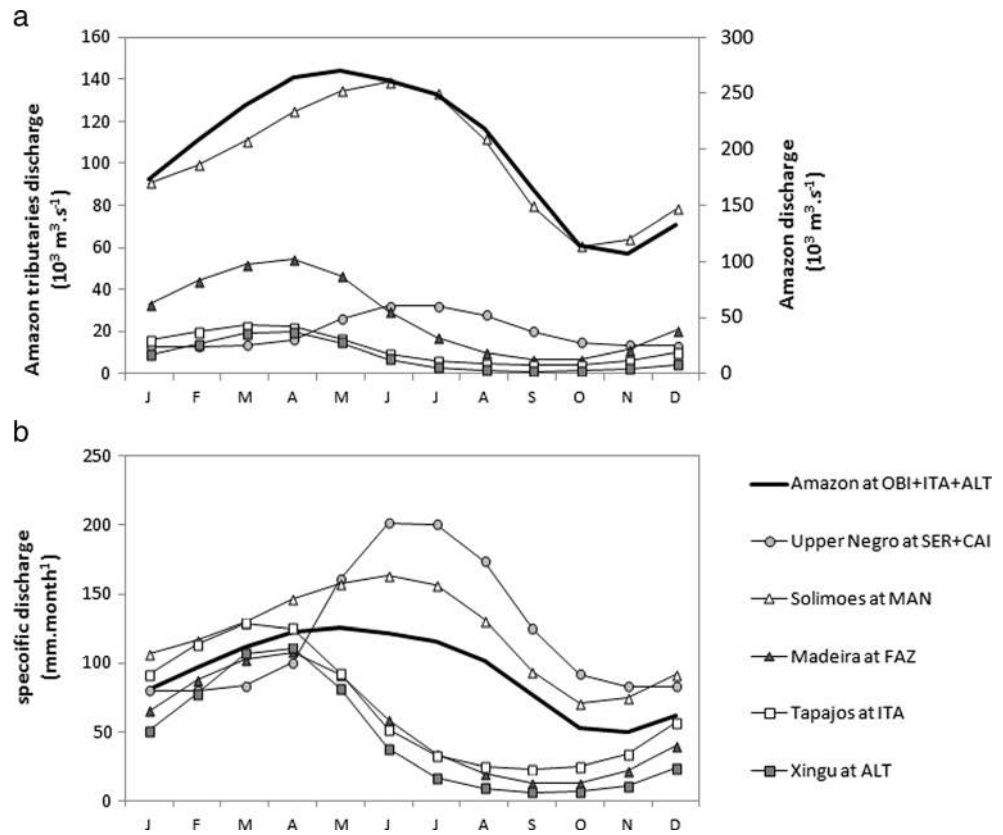
(Fig. 2b). Indeed, the Tapajós River experiences the highest seasonality and the lowest annual specific discharge, whereas the Negro River has the lowest seasonality but the highest specific discharge. The Amazon River at the Óbidos station displays the highest discharge between April and June. This maximum in discharge reflects the inputs of the Solimões River, but the seasonal variability of discharge of the Amazon at the Óbidos station is smoothed by the lag in time of the maximum discharge in the tributaries (the Madeira, Solimões and Negro Rivers) over the years (Fig. 2) and by the storage in the floodplains (Guimberteau et al. 2012).

The mean transit times of overland flow, shallow subsurface flow and groundwater have been estimated at the scale of the main Amazon tributaries based on the hydrograph separation using water-stable isotopes (Mortatti et al. 1997; Tardy et al. 2005; Bustillo et al. 2011). For example, Tardy et al. (2005) showed that the mean half-lives of water in the Amazon basin are approximately 22.2, 32.6 and 60.5 days for superficial runoff, hypodermic and groundwater reservoirs which annually contribute 29.5, 47.8 and 22.7 % of the Amazon discharge, respectively. Recently, based on hydrological models, Paiva et al. (2013) showed that surface waters govern most of the monthly terrestrial water storage changes (56 %), followed by soil water (27 %) and ground water (8 %). In the channel, as the velocity of surface water ranges between 1 and 2 m s⁻¹ (SO-HYBAM gauging database), the mean transit time from the Andes to the outlet of the basin (approximately 6,000 km) can be higher than 1 month.

Material and methods

In this study, we used the available data of the Programa Climatológico y Hidrológico de la Cuenca Amazonica de Bolivia (PHICAB) and HYBAM observatory (Geodynamical, hydrological and biogeochemical control of erosion/alteration and material transport in the Amazon, Orinoco and Congo basins—Environmental Research Observatory) frameworks (the HYBAM data are freely available at the HYBAM observatory website, <http://www.ore-hybam.org>, and the PHICAB database is available in Guyot 1993). The PHICAB framework collected hydrochemical data at various gauging stations of the Upper Madeira basin between 1983 and 1992. The HYBAM framework has collected hydrochemical data since 2000. In the present study, we use the available data corresponding to the 2000–2012 period in the Solimões basin, at three stations in the Madeira basin and at Brazilian stations on the Negro, Amazon and Tapajós Rivers (Table 2). Importantly, the studied stations are not homogeneously distributed throughout the Amazon basin. Among the 34 studied stations, 26 are located at the outlet of the Andean and foreland basins, seven stations record the inputs of the main Amazon tributaries, and Óbidos (OBI) is

Fig. 2 **a** Monthly discharge of the Amazon River (*right axis*) and its main tributaries (*left axis*) and **b** monthly specific discharge of these rivers. Data are averaged for the 1983–2013 period (source: SO-HYBAM website)



the main station recording the Amazon outputs. Each gauging station is hereafter named according to a three-letter code listed in Table 2.

Hydrological data

Water levels were collected daily or twice daily using a conventional hydrologic method. During the PHICAB period (1983–1992), gauging was accomplished using punctual velocity measurements with a mechanical current metre over the whole river section (Guyot 1993). For the 2000–2012 period, within the framework of the HYBAM observatory (<http://www.ore-hybam.org>), gauging was accomplished using a 600-kHz (Peru, Bolivia and Brazil) or 1,200-kHz (Ecuador) acoustic Doppler current profiler (ADCP). The daily discharge series was then calculated from rating curves (discharge–water level relationship) using the Hydraccess software (Vauchel 2005; free download at the SO-HYBAM website).

Sampling method and water chemistry analysis

Conductivity and pH were measured every 10 days at a set of stations of the HYBAM framework, named the ‘SO-HYBAM framework’ and following a 3-month frequency at the other stations of the HYBAM network (named the ‘reference HYBAM network’) as well as at the PHICAB stations.

Remarkably, a conductivity time series is available at a frequency of 10 days since October 1994 at the Óbidos station.

Regarding the hydrochemical data, a sample of 650 ml was collected monthly at each gauging station, with the exception of the ALT station (Xingu River), where we use only two sample analyses. The samples were filtered on site using a 0.45- or 0.2- μm porosity filter. Major element concentrations (Ca, Mg, Na, K, Cl, SO_4 , HCO_3 and Si) were determined at the Instituto de Investigacion en Quimica, Universidad Mayor de San Andres (IIQ-UMSA) laboratory (La Paz, Bolivia) for the 1983–1992 sampling period, at the Géoscience Environnement Toulouse laboratory (GET, France) for the 2000–2012 sampling period for Ecuadorian, Peruvian and Bolivian samples and at the Laboratory of Geochronology, Universidade de Brasilia (UnB) for the 2003–2012 sampling period for Brazilian samples. The samples collected during the PHICAB program were analysed according to the analytical procedure described in Guyot (1993): alkalinity was measured by acid titration, Cl^- by mercuric thiocyanate colorimetry, SO_4^{2-} by barium sulfate nephelometry, Ca^{2+} and Mg^{2+} by atomic absorption, Na^+ and K^+ by flame spectrometry and Si by ammonium molybdate colourimetry. The samples collected during the HYBAM program were analysed according to the procedure described in Cochonneau et al. (2006). Cl^- and SO_4^{2-} concentrations were measured by ion chromatography, and Na^+ , Ca^{2+} , Mg^{2+} , K^+ and Si were analysed by inductively coupled plasma-atomic emission spectroscopy

(ICP-AES). Alkalinity was determined by acid titration. The TDS concentration corresponds to the sum of the cations (Ca, Mg, Na and K), the anions (HCO_3^- , SO_4^{2-} and Cl) and SiO_2 concentrations, all expressed in milligram per litre. Based on the analyses of geostandards, the analytical error is less than 10 % for both sampling periods.

We observed that TDS and conductivity are significantly correlated ($r^2=0.86$; p values <0.01 ; $N=1,782$), considering the whole database, with the exception of one outlier (when conductivity or TDS $>$ average $+3\sigma$; ABA station):

$$\text{TDS}(\text{mg L}^{-1}) = 0.8 \times \text{conductivity}(\mu\text{S cm}^{-1}) \pm 21 \text{ mg L}^{-1}$$

Here, the error is defined as the mean absolute error (MAE) of the relationship.

In the present study, we use the conductivity as a proxy for TDS concentration to describe the concentration (C) vs discharge (Q) relationship because of the higher frequency of conductivity acquisition (10 days), allowing a precise characterization of the total solute dynamics throughout the hydrological year.

Annual flux calculation

The Amazon TDS flux corresponds to the sum of the inputs of the Amazon River at the OBI station, the Tapajós River at the ITA station and the Xingu River at the ALT station. To calculate the annual flux of each solute at each station, we first estimated the monthly flux based on the multiplication of the instantaneous concentrations obtained at a monthly period by the monthly discharge calculated from the sum of daily discharges (method M1C in Table S1 of the supplementary material). We then calculated the interannual average of monthly fluxes. We finally added the 12 monthly flux averages to evaluate the mean annual flux.

At two stations (the OBI and ITA stations), we also performed three other annual flux calculations for TDS (Table S1) to compare the effect of the flux calculation methods based on the same dataset in the period 2003–2012. The results are presented in Table S2. The method M1A corresponds to the mean concentration of the overall data multiplied by the mean monthly discharge; then, we added the 12 monthly flux averages to evaluate the mean annual flux. The method M1B corresponds to the monthly concentration interannual average multiplied by the mean monthly discharge interannual average; then, we added the 12 monthly flux averages to evaluate the mean annual flux. The method M2A corresponds to the application of the $C=aQ^b$ relationship (see the following section) based on the daily discharge database. The daily flux is then calculated based on the multiplication of C_j and Q_j , and the annual flux corresponds to the sum of the daily fluxes.

The error associated with these estimates corresponds to the sum of the analytical error ($<10\%$) and the error associated with the discharge measurements. The discharge measurement errors at OBI, ITA and ALT were estimated to be ± 3 , ± 10.7 and $\pm 9.8\%$, respectively, by Callède et al. (2010). The weighted discharge measurement error of the sum of the discharge at these three stations was therefore $\pm 4\%$. A maximum of error of 14 % is estimated. Indeed, regardless of the calculation method performed, if we compare the sum of the annual solute fluxes calculated upstream of the OBI station (at the MAN, FAZ, CAI and SER stations) with the solute fluxes calculated at the OBI station, the upstream flux is equal to the OBI fluxes within 10 % ((flux OBI–flux upstream)/flux OBI) for TDS, Cl, SO_4 , HCO_3^- , Ca, Mg and Na. Only the upstream annual fluxes of K and Si are 11 and 26 % lower than those at the flux measured at OBI station most likely due to significant K and Si inputs by the surface basin included between these stations.

C vs Q relationships

Water discharge is an important factor affecting the variation of riverine material fluxes. The time series of dissolved concentrations allows us to derive the relationships between major element concentrations and river discharge. These relationships are functionally described using:

$$C_i = a Q_i^b \quad \text{or} \quad \log(C_i) = 10^a + b \log(Q_i) \quad (1)$$

where C_i is the instantaneous (daily) concentration of the studied major element, Q_i is the daily discharge corresponding to the sampling date and a and b are the regression coefficients. Such relationships have been reported for several rivers to characterize the variation of the solute concentration in response to discharge variability (e.g. Walling and Webb 1983; Nkounkou and Probst 1987; Roy et al. 1999; Chen et al. 2002; Godsey et al. 2009; Gislason et al. 2009; Clow and Mast 2010; Li et al. 2012; Laraque et al. 2013; Moon et al. 2014). These relationships between flow and load have important implications for predicting the fluxes of chemical constituents to receiving water bodies due to variable environmental conditions (Godsey et al. 2009; Jawitz and Mitchell 2011). In this paper, we used the slope b of the C – Q regression as an index, allowing for the characterization of the first-order response of the production of Amazonian solutes to changes in discharge. Indeed, the slope is a measure of the first-order degree to which concentration responds to a change in discharge. The larger the slope, the less sensitive is the concentration. When $b \sim 0$, the concentration variability is low and independent of discharge. This behaviour, called ‘chemostatic’ (Godsey et al. 2009), reflects a constant solute concentration despite variations in discharge. When $b \sim -1$, concentration decreases with increasing discharge, corresponding to a total dilution pattern and reflecting a constant

solute flux despite variations in discharge. When $b > 0$, concentration increases with increasing discharge, as generally observed for suspended load concentrations (e.g. Armijos et al. 2013b).

Based on a global compilation that included HYBAM for the Amazon at the OBI station, Moon et al. (2014) suggested that at least 40 paired concentration–discharge measurements are needed to calculate unbiased weathering rates. Following this criterion, the data from the PUN, PCH and ALT stations are excluded for the analysis. Moreover, the C – Q relationship at NYO station is not significant (Moquet et al. 2014b). In the supplementary material S3, we reported the regression parameters with their related standard errors (SE) on a and b values and the mean absolute percentage error (MAE%) of each regression. Because of the occurrence of hysteresis behaviour at the OBI station (see below), we did not use this method to calculate the annual Amazon fluxes.

Results

Hydrochemical characteristics

The description presented hereafter is based on the PHICAB and HYBAM monitoring networks and partly summarizes results already reported in Moquet et al. (2011), for the Andean tributaries, and in Sanchez et al. (2015), for some of the downstream stations. Even if slight differences in the mean concentration values are observed between the present study and the previous ones due to the actualization of the database, the values measured in both studies are commensurate. Hydrochemical characteristics are reported in Table 3 and are illustrated in Fig. 3.

The pH of the river water samples ranges between 4.8 and 7.9 (median value for each gauging stations). The highest value is recorded in the semi-arid basin of the Grande River (ABA station, Upper Madeira River), which is characterized by significant carbonates and Neogene evaporites (Moquet et al. 2011). The lowest pH values are recorded in the Upper Negro River at the SER station, which is characterized by a high abundance of arenosol/podzol soils and is known to release waters with high organic matter concentrations (e.g. Stallard and Edmond 1983; Moreira-Turcq et al. 2003). At the other stations, the pH is closer to neutral conditions, with slightly alkaline water in the Andean tributaries and slightly acid water in the rivers of the central plain and shield. These values are coherent with previous observations (Sioli 1964; Junk and Piedade 1997; Rios-Villamizar et al. 2014).

The TDS concentration (and conductivity) is contrasted throughout the basin (Fig. 3 and Table 3). The highest mean value (TDS at ABA=396 mg L⁻¹) is recorded in the semi-arid basin of the Grande River (Upper Madeira), and the lowest value (TDS at SER=6.5 mg L⁻¹) is recorded in the podzolic/arenitic basin of the Upper Negro River. The other TDS values

decrease as follows: Marañon–Ucayali (TDS=63 to 222 mg L⁻¹) > Napo (TDS=64 to 99 mg L⁻¹) = Upper Madeira (TDS=63 to 109 mg L⁻¹) > central plain tributaries (TDS of Purus at LAB=52 mg L⁻¹) > shield tributaries (Negro, Branco, Tapajós and Xingu Rivers; TDS=22 to 48 mg L⁻¹). This hierarchy reflects the lithological/geomorphological characteristics of the river basins. Importantly, this hierarchy is based only on the monitored tributaries that integrate the upstream sub-basins. Intra-basin concentration heterogeneities have previously been observed, especially in the Andes (e.g. Stallard and Edmond 1983; Guyot et al. 1998). The Grande River (Madeira Andes) is characterized by significant evaporite deposits and a semi-arid climate. The Marañon and the Ucayali (Andean tributaries) drain large evaporite and carbonate outcrops. The Napo basin (Andean tributaries) includes the main active volcanoes of the basin. The central plain tributaries (e.g. Purus River) drain old deposits of silicate sediments, initially exported from the Andean area. These waters are fairly diluted. As a consequence, the Solimões (TAM, TAB and MAN) and Madeira (PVE and FAZ) stations are affected by dilution associated with the foreland and central plain inputs. The tributaries in the shields (Negro, Branco, Tapajós and Xingu Rivers at the SER, CAI, ITA and ALT stations, respectively) are characterized by highly diluted waters. Even if all the basins of the shields exhibit low TDS concentrations (<35 mg L⁻¹), they can be separated into two groups: the Branco, Tapajós and Xingu Rivers (CAI, ITA and ALT stations, respectively; 22–35 mg L⁻¹) exhibit higher TDS concentrations than the Upper Negro River (SER station; 6 mg L⁻¹).

With the exception of some specific contexts, Ca²⁺, Mg²⁺, Na⁺, SO₄²⁻, Cl⁻ and HCO₃⁻ concentrations follow the same hierarchy as the conductivity and TDS. However, high relative chlorine and sodium contents are recorded in the Marañon sub-basin in the Huallaga River (CHA station), which drains large evaporitic domes (Benavides 1968) and in the El Tigre River (NYO station), which was characterized by formation water released from oil extraction activity until February 2009 (Moquet et al. 2014b) (Fig. 3). These two basins exhibit high Cl and Na concentrations (Cl⁻ and Na⁺ ≥800 μmol L⁻¹) and a 1:1 Cl/Na ratio characteristic of halite dissolution. SO₄²⁻ concentrations are generally higher than 50 μmol L⁻¹ in all Andean sub-basins. This element is released by dissolution of evaporites (gypsum) and by pyrite oxidation (Berner and Berner 1987; Moquet et al. 2011).

In contrast with TDS and the Ca²⁺, Mg²⁺, Na⁺, SO₄²⁻, Cl⁻ and HCO₃⁻ concentrations, K⁺ and Si do not follow the same trend. The highest concentrations of dissolved Si are measured in the basins draining volcanic areas (the Upper Napo and the PUN and SAN stations of the Marañon basin), and the lowest values are measured in the Upper Negro River. Despite the high geographical diversity (lithology, geomorphology, climate, etc.), all the other basins exhibit a low spatial variability of dissolved Si concentration. High K⁺ concentrations are measured in the El Grande River (ABA station), probably

Table 3 Physico-chemical characteristics and average dissolved solid concentrations for the monitored stations (for the ALT station, the values for the two analysed samples are presented)

Station name	pH	Cond $\mu\text{S cm}^{-1}$	Ca $\mu\text{mol L}^{-1}$	Mg	Na	K	Cl	SO ₄	HCO ₃	SiO ₂	TDS mg L^{-1}
	Median (min–max)	Avg $\pm\sigma$									
NLO	7.1 (6–8.8)	92 \pm 16	318 \pm 55	57 \pm 11	98 \pm 30	38 \pm 38	31 \pm 64	48 \pm 13	724 \pm 140	208 \pm 42	80 \pm 13
DJI	7.2 (5–8.7)	88 \pm 31	233 \pm 133	52 \pm 22	114 \pm 47	41 \pm 34	25 \pm 19	49 \pm 31	648 \pm 253	231 \pm 79	77 \pm 24
FOR	7 (6.3–7.8)	63 \pm 19	189 \pm 41	49 \pm 13	116 \pm 48	36 \pm 18	20 \pm 12	29 \pm 10	544 \pm 123	251 \pm 46	64 \pm 10
SEB	7.2 (6.5–8.1)	118 \pm 19	339 \pm 68	110 \pm 32	172 \pm 49	42 \pm 15	66 \pm 44	110 \pm 42	781 \pm 129	289 \pm 54	98 \pm 15
NRO	6.8 (6.2–7.5)	71 \pm 24	192 \pm 46	68 \pm 42	133 \pm 59	36 \pm 7	29 \pm 18	42 \pm 18	566 \pm 262	278 \pm 98	71 \pm 25
BEL	6 (5.1–7.3)	43 \pm 12	129 \pm 46	48 \pm 17	93 \pm 65	26 \pm 13	43 \pm 79	25 \pm 13	404 \pm 114	205 \pm 39	50 \pm 10
SAN	7.4 (6.3–9.6)	71 \pm 13	207 \pm 45	60 \pm 11	118 \pm 31	23 \pm 8	27 \pm 8	50 \pm 12	526 \pm 135	200 \pm 52	63 \pm 10
BOR	7.1 (5.9–8.2)	172 \pm 56	615 \pm 249	99 \pm 28	217 \pm 100	27 \pm 6	128 \pm 144	110 \pm 78	1,337 \pm 476	192 \pm 61	140 \pm 42
PUN	7.2 (5.4–8.8)	111 \pm 29	215 \pm 58	181 \pm 67	308 \pm 107	43 \pm 12	74 \pm 29	105 \pm 37	826 \pm 247	260 \pm 89	100 \pm 27
CHA	6.8 (6–8)	292 \pm 76	784 \pm 126	120 \pm 45	1,054 \pm 745	33 \pm 9	949 \pm 745	181 \pm 99	1,595 \pm 266	180 \pm 25	222 \pm 52
NYO	5.7 (4.8–7.5)	138 \pm 95	144 \pm 146	48 \pm 36	788 \pm 543	21 \pm 7	897 \pm 731	28 \pm 48	378 \pm 389	170 \pm 45	103 \pm 62
SRE	6.8 (5.4–9.6)	160 \pm 54	489 \pm 144	83 \pm 25	532 \pm 290	27 \pm 6	532 \pm 432	72 \pm 40	1,100 \pm 281	190 \pm 43	145 \pm 33
ATA	7.4 (5.9–8.2)	254 \pm 68	792 \pm 209	179 \pm 57	490 \pm 258	44 \pm 9	334 \pm 203	294 \pm 129	1,571 \pm 361	211 \pm 51	196 \pm 42
JHE	6.6 (5.6–7.4)	209 \pm 112	713 \pm 327	129 \pm 62	386 \pm 262	37 \pm 10	282 \pm 225	153 \pm 93	1,440 \pm 757	161 \pm 41	155 \pm 80
TAM	6.7 (5.8–8)	194 \pm 47	652 \pm 101	113 \pm 22	446 \pm 180	34 \pm 9	365 \pm 167	116 \pm 29	1,358 \pm 224	182 \pm 33	158 \pm 23
TAB	7.4 (6.2–8.1)	147 \pm 23	508 \pm 101	99 \pm 22	343 \pm 136	33 \pm 5	231 \pm 99	87 \pm 24	1,082 \pm 207	211 \pm 72	128 \pm 16
LAB	6.9 (5.9–7.7)	42 \pm 22	149 \pm 79	51 \pm 26	78 \pm 41	28 \pm 8	16 \pm 14	25 \pm 15	355 \pm 209	230 \pm 86	51 \pm 18
MAN	6.9 (4–7.8)	77 \pm 18	254 \pm 64	57 \pm 12	159 \pm 60	25 \pm 7	110 \pm 50	45 \pm 20	593 \pm 135	158 \pm 23	70 \pm 12
VGR	6.7 (3.6–9.6)	40 \pm 20	63 \pm 73	47 \pm 34	109 \pm 68	51 \pm 27	65 \pm 76	29 \pm 59	386 \pm 163	174 \pm 71	47 \pm 16
ABA	7.9 (7–9)	625 \pm 164	1,091 \pm 655	1,052 \pm 636	1,435 \pm 666	119 \pm 45	701 \pm 629	1,149 \pm 937	2,344 \pm 489	155 \pm 62	396 \pm 131
PVI	6.6 (3.8–10)	83 \pm 24	141 \pm 107	89 \pm 43	202 \pm 199	40 \pm 48	178 \pm 545	160 \pm 108	273 \pm 197	169 \pm 84	63 \pm 33
PSI	7.6 (6.4–9.7)	142 \pm 51	257 \pm 103	199 \pm 83	289 \pm 181	60 \pm 19	138 \pm 144	144 \pm 124	878 \pm 388	141 \pm 49	109 \pm 41
GUA	7.6 (5.7–9.6)	91 \pm 32	171 \pm 73	118 \pm 52	168 \pm 92	50 \pm 10	72 \pm 57	93 \pm 76	575 \pm 174	160 \pm 79	73 \pm 21
RUR	7 (6.2–8.4)	128 \pm 13	270 \pm 66	169 \pm 49	164 \pm 57	29 \pm 9	30 \pm 16	264 \pm 85	556 \pm 123	146 \pm 22	89 \pm 20
PCH	7.2 (5.1–8.8)	122 \pm 24	264 \pm 83	174 \pm 79	148 \pm 69	43 \pm 13	66 \pm 60	130 \pm 95	730 \pm 148	157 \pm 71	89 \pm 16
MIR	7.2 (4–10.3)	77 \pm 28	197 \pm 82	78 \pm 42	125 \pm 70	40 \pm 15	86 \pm 98	60 \pm 43	709 \pm 337	181 \pm 54	79 \pm 28
CES	7.1 (4.7–10)	86 \pm 22	194 \pm 63	103 \pm 42	119 \pm 57	40 \pm 12	75 \pm 81	83 \pm 56	630 \pm 156	173 \pm 65	75 \pm 15
PVE	7.2 (5.8–8)	77 \pm 22	214 \pm 76	113 \pm 36	126 \pm 51	38 \pm 6	28 \pm 21	101 \pm 55	534 \pm 158	178 \pm 64	69 \pm 17
FAZ	6.9 (5.9–7.6)	53 \pm 17	139 \pm 46	81 \pm 25	112 \pm 58	36 \pm 8	26 \pm 16	57 \pm 40	381 \pm 112	150 \pm 39	50 \pm 11
SER	4.8 (3.6–6.7)	7 \pm 4	10 \pm 5	5 \pm 2	26 \pm 37	10 \pm 7	7 \pm 4	2 \pm 5	18 \pm 19	65 \pm 12	6 \pm 1
CAI	6.6 (5.2–7.6)	27 \pm 9	41 \pm 9	34 \pm 7	98 \pm 41	36 \pm 14	22 \pm 11	7 \pm 9	224 \pm 77	229 \pm 38	35 \pm 7
OBI	6.9 (6.2–7.5)	51 \pm 12	151 \pm 43	44 \pm 13	97 \pm 36	23 \pm 5	58 \pm 26	32 \pm 15	381 \pm 102	163 \pm 49	48 \pm 11
ITA	6.5 (5.3–7.5)	16 \pm 4	32 \pm 17	23 \pm 5	32 \pm 11	22 \pm 4	17 \pm 14	2 \pm 2	143 \pm 52	153 \pm 21	22 \pm 4
ALT	(6.7–6.6)	14–17	31–32	24–23	35–64	19–25	14–15	2–2	174–192	118–153	22–26

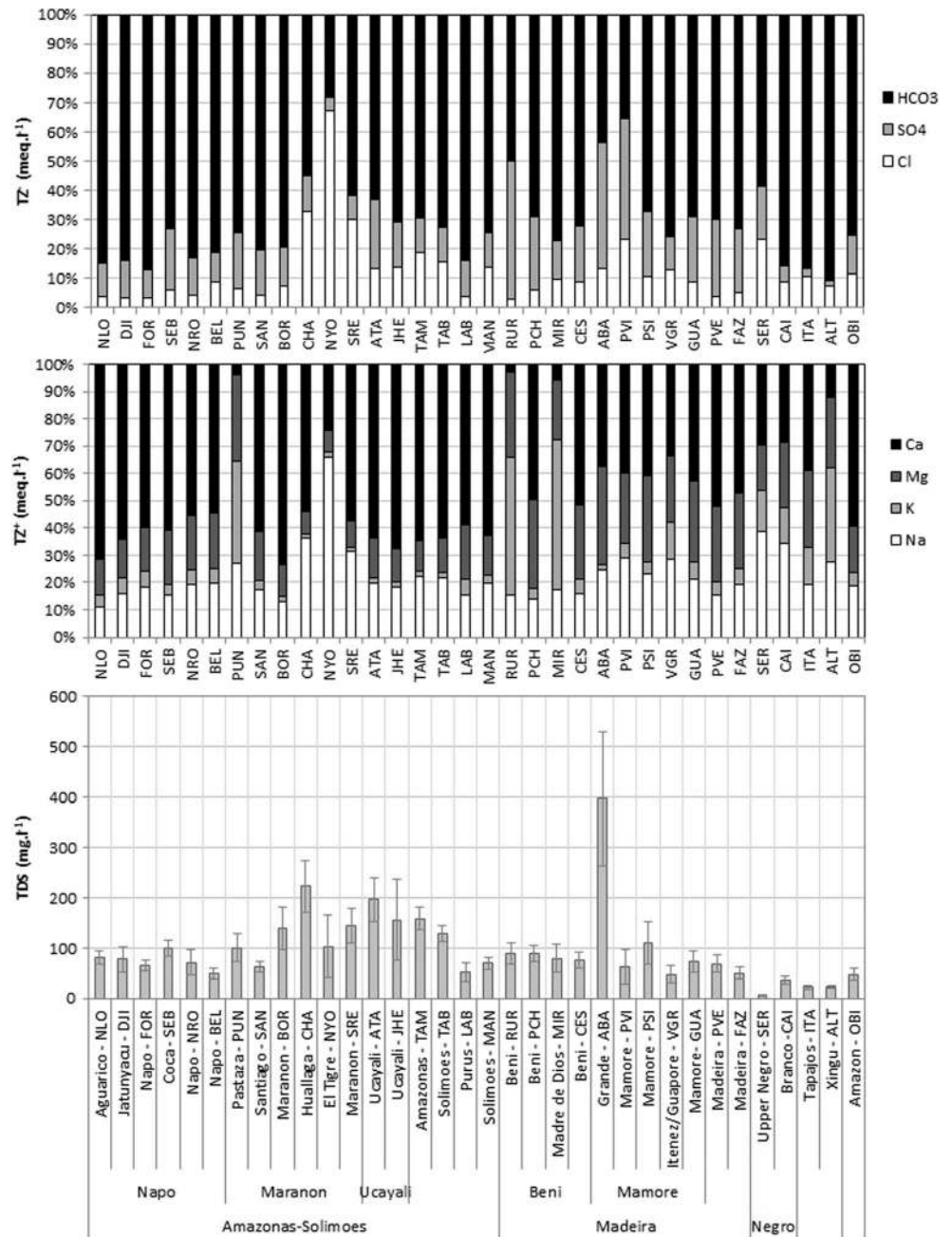
due to the high inputs of evaporite dissolution (Guyot 1993). In the other rivers, the K⁺ concentration has low spatial variability. These elements can both reflect silicate weathering and vegetation inputs (Berner and Berner 1987; Lucas 2001; Chaudhuri et al. 2007). With the exception of some specific contexts (volcanic area and upper Negro River for Si and Grande River for K⁺), the Si and K⁺ concentrations are relatively homogenous over areas characterized by high heterogeneity in lithology, geomorphology or climate. This difference

in the spatial variability in concentration between K⁺ and Si and the other major elements was also observed by Guyot et al. (1998) in the Madeira basin.

C–Q relationships

Regression parameters of Eq. 1 were determined for TDS concentration, conductivity and all elements over all gauging stations (Fig. 4).

Fig. 3 Dissolved load chemical composition and average conductivity values (\pm standard deviation) of the monitored rivers



The slope b of the $C-Q$ relationship varies from station to station and from element to element, but b value is usually between 0 and -1 in the Amazon basin, indicating that concentrations remain nearly constant or respond inversely to changes in discharge, consistently with observations at a global scale (Meybeck et al. 1996).

The definition of a $C-Q$ log relationship is not adapted for all elements in all contexts to reproduce the measured concentration. A MAE greater than 50 % is generally measured for Cl and/or SO₄ in the Solimões upper basin at the SRE and BEL stations and in the upper Madeira basin and for most of the elements at the SER station, indicating that the observed $C-Q$ relationships are non-linear in log-log space (Fig. 4). At the

SER station, this observation can be explained by the very low concentrations and high relative error. At the other stations with high MAE, the $C-Q$ relationships are simply not univocal.

As TDS and conductivity are correlated, they generally exhibit similar b values under the ± 1 SE uncertainties (Figs. 4 and 5). Throughout the basin, the TDS b value varies between -0.42 ± 0.07 and 0.02 ± 0.04 but is generally close to -0.20 . This implies that the TDS variability is lower than the discharge variability and that, at first order and as observed in other contexts (e.g. Markewitz et al. 2006; Yuan et al. 2007), the TDS flux variability is controlled by discharge along the hydrological cycle.

Figure 5 shows conductivity vs discharge at nine selected stations representative of the main geomorphological contexts of the Amazon basin. The Solimões River (BOR, TAB and MAN stations) exhibits the highest conductivity values. In its Andean part (BOR station), the conductivity is highly variable. This behaviour can reflect the spatial and temporal heterogeneity of rainfall and solute sources. Downstream at the TAB and MAN stations, lower conductivity values can reflect the dilution effect due to the contribution of tributaries draining sedimentary areas. Nevertheless, in response to the hydrological cycle at all these stations, the variability of conductivity remains low in comparison with discharge variability, and b value is lower than -0.2 .

In the Madeira sub-basins, the conductivity– Q slope b is nearly zero at the RUR station, representing the Andean part of the sub-basin, and decreases to -0.2 in the central plain. At the RUR station, even if the slope b is close to zero, concentration is variable, as previously observed by Guyot et al. (1993), and two regimes can be distinguished. For discharge lower than $1,800 \text{ m}^3 \text{ s}^{-1}$, the conductivity is close to the simple dilution curve. For discharge higher than $1,800 \text{ m}^3 \text{ s}^{-1}$, conductivity is more variable and in some cases increases with discharge. The hypothesis considered here is that, as observed in other Amazonian contexts (Markewitz et al. 2001), the highest concentration during the rainy period (November–May), when discharge is higher than $1,800 \text{ m}^3 \text{ s}^{-1}$, can reflect inputs from the mobilization of elements by surface runoff from the upper layer of the soils. Such a contribution might vary in the Andes from one rain event to another due to the spatial and temporal rainfall heterogeneity (Roche et al. 1990; Guyot et al. 1993). During the dry period, the variability in concentration would be mainly controlled by the dilution effect associated with groundwater inputs. Downstream, on the Madeira basins at the PVE and FAZ stations, the conductivity is less variable for a given discharge range. The conductivity decreases with discharge, but b value remains high ($b \sim -0.2$) and the flux variability is therefore mainly controlled by discharge variability. This dilution effect has been previously reported by Roche and Fernandez Jauregui (1988) in plain basins of the Madeira.

In the basins of the shields (SER, CAI and ITA), the conductivity remains relatively constant (with b value higher than -0.1 and lower than 0.1) in response to the hydrological cycle.

The b values for Ca, Mg, SO_4 and HCO_3 are higher than -0.5 but are generally lower than 0 . For Ca, Mg, HCO_3 and TDS, the Solimões stations exhibit generally higher b values than the Madeira stations, as previously shown by Sanchez et al. (2015). This implies that the Solimões basins are more responsive to rainfall events in term of solute delivery than the Madeira basins, which tend to dilute to a limited extent. Cl and Na exhibit very variable b values ranging from -0.78 ± 0.09 to 0.04 ± 0.06 , and the lowest b values are generally recorded at

some Napo stations, Marañon-Ucayali stations and downstream Solimões and Amazon stations (TAB, MAN and OBI), whereas b values closer to 0 are recorded in the plain (LAB) and in the tributaries in the shields (CAI and ITA). The Napo River exhibits an overall decrease of b from the Andean stations to the foreland station (BEL). This implies that the foreland area of this basin contributes more to diluting the Andean inputs than it contributes to solute inputs. Among the solutes, the b values of Si and K are close to 0 in most cases. With the exception of Cl and Na, these results imply that, at first order, discharge variability is the main factor controlling the variability of solute and TDS fluxes throughout the year.

These C – Q regressions must be considered as indexes that allow characterizing the first-order relationship between the variability of solutes and their respective daily discharge using a large amount of data from 30 stations of the Amazon basin. In specific cases, these relationships are not significant (e.g. NYO station, Moquet et al. 2014b), and in other cases, other functional relationships can better explain the temporal variability in concentration. This is the case in particular when a significant hysteresis is observed. We illustrate this latter case through the description of hysteresis behaviour observed at the OBI station.

At the OBI station, the C – Q regressions defined for conductivity and TDS concentration allow estimating the measured values with a mean average error of $\pm 8 \mu\text{S cm}^{-1}$ and $\pm 7.5 \text{ mg L}^{-1}$, respectively. Nevertheless, examining closely the TDS and the conductivity as a function of discharge, a non-univocal C – Q relationship is observed (Fig. 6a). The relationship between conductivity and discharge of the Amazon at the OBI station follows a clockwise hysteresis and concave curvature behaviour ('C1' type hysteresis of Evans and Davies 1998 classification) (Fig. 6a). The highest conductivity values ($>70 \mu\text{S cm}^{-1}$) are recorded during the low water stage (November–December), whereas lower values are recorded during the high water stage (June–August period). For a given discharge value, higher conductivities associated with the increasing water stage (January–May) compared with the decreasing water stage (September–October) are recorded. Here, this absence of a univocal relationship between TDS and discharge is clearly due to a mass water mixing phenomenon that is the combined effect of (1) the contrast of the mean TDS concentration/conductivity between the three main tributaries (Negro, Solimões and Madeira Rivers) and (2) the lag of their relative discharge contribution over the course of the year. The monthly TDS flux at the OBI station corresponds to the sum of the inputs of the Solimões River at the MAN station, the Madeira River at the FAZ station and the Negro River at the CAI and SER stations within a range of $\pm 11 \%$ (relative difference between the monthly OBI fluxes and the upstream tributary fluxes) (Fig. 6b). Following this calculation, the Solimões controls

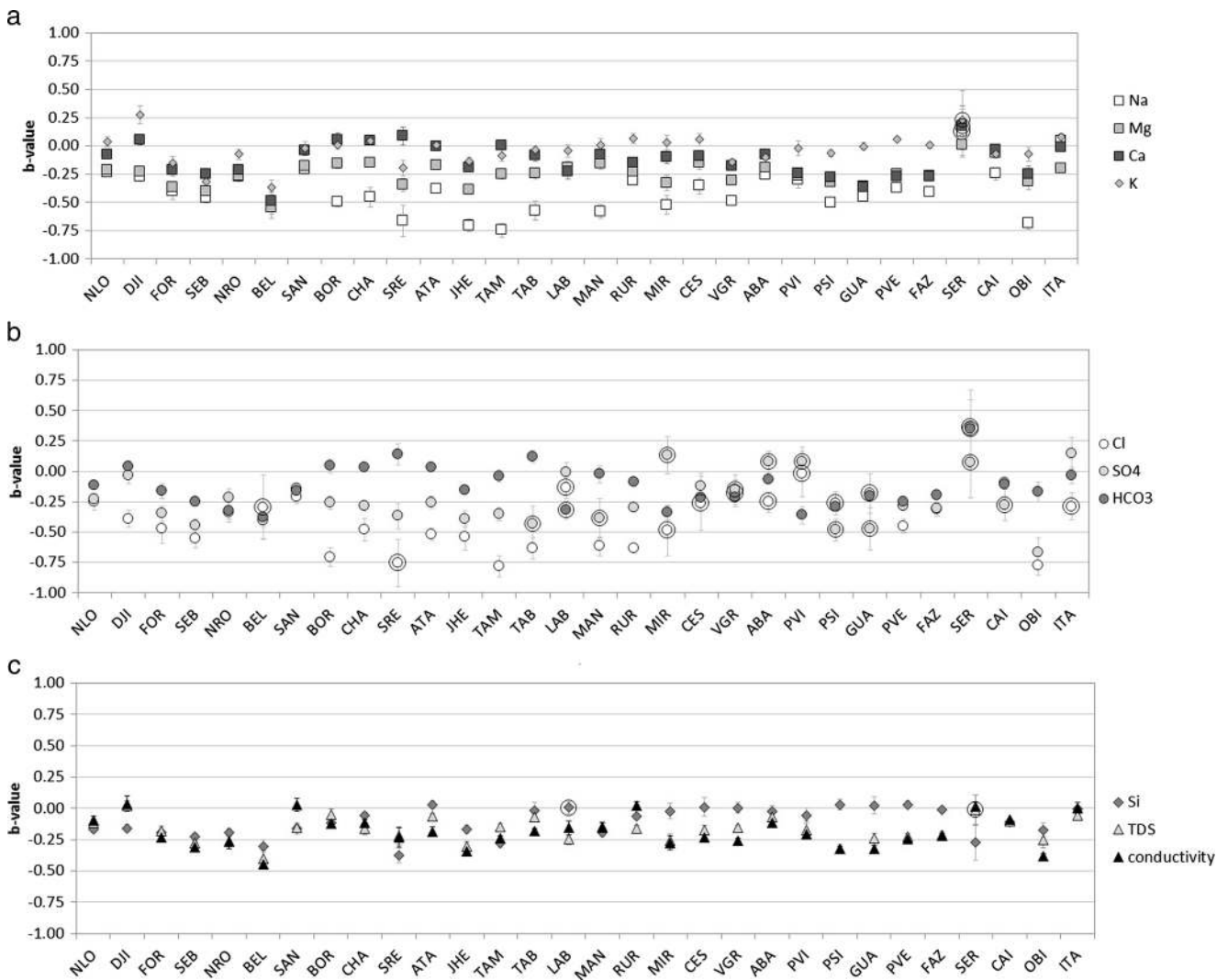


Fig. 4 Log-log slopes of concentration–discharge relationships ($\log(C_i)=a+b \log(Q_i)$) for studied stations by site name for cations (a), anions (b), TDS, Si and conductivity (c). Error bars indicate ± 1 standard error (SE). Large symbols indicate that the MAE of the C–Q relationship is $> 50\%$

between 77 and 89 % of the monthly TDS inputs in February and September, respectively. In contrast, the Madeira River controls between 8 and 22 % of the TDS inputs in September and February, respectively. The Negro River inputs at the CAI and SER stations control less than 5 % of the monthly TDS flux over the course of the year. The hysteresis behaviour is also explained by the variation in the relative discharge contribution of the Solimões, the Madeira and the Negro Rivers (Fig. 2). During the November–January period, when TDS concentration is at a maximum at the Óbidos station, the Solimões at the MAN station controls more than 60 % of the Amazon discharge at the OBI station. During the February–June period, the relative contribution of the Solimões discharge decreases as the discharge of the Madeira continues to contribute more than 20 % of the Amazon discharge. The TDS flux of the Solimões River is consequently partly diluted by that of the Madeira River. During the July–October period, the

discharge inputs of the Negro River and the discharge produced by the remaining area between the Negro, Solimões and Madeira River stations and the OBI station are maximal and represent more than 30 % of the Óbidos discharge. These highly diluted water inputs would consequently dilute the mineralized Solimões and Madeira River water inputs. To confirm this behaviour, we calculated the TDS concentration at the OBI station following Eq. 2:

$$C_{OBI} = (F_{MAN} + F_{FAZ} + F_{SER} + F_{CAI})/Q_{OBI} \quad (2)$$

where C_{OBI} is the mean monthly concentration (mg L^{-1}), F is the mean monthly flux at the MAN, FAZ, SER and CAI stations (mg month^{-1}) and Q_{OBI} is the mean monthly discharge at the OBI station (L month^{-1}).

This calculation reproduces the TDS concentration at the OBI station within a range of $\pm 6 \text{ mg L}^{-1}$ (Fig. 6c).

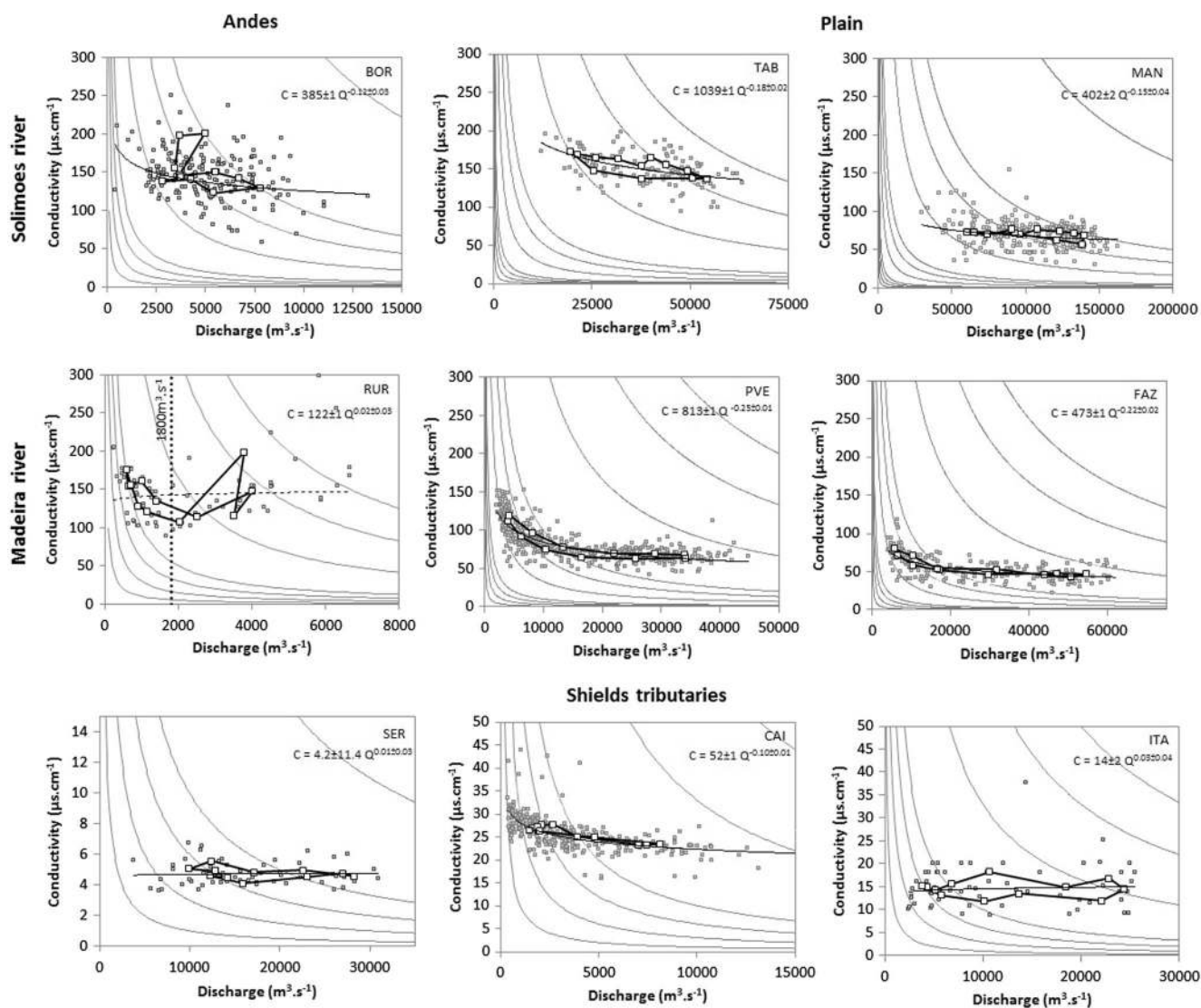


Fig. 5 Ten-day frequency conductivity measurements (*small symbols*) and monthly averages (*large symbols and lines*) plotted against discharge for nine selected gauging stations. Simple dilution curves (concentration variability of a constant flux) are added for reference

Not all individual solutes show hysteresis. As Ca and HCO_3 are the main species contributing to the TDS content (Fig. 3), they control the hysteresis behaviour observed for TDS and conductivity and also individually show hysteresis (Fig. S1). Hysteresis phenomena are also observed for Mg, SO_4 and K. Cl and Na exhibit dilution behaviour because these elements are mainly produced by the Solimões basin. Consequently, the Negro and Madeira Rivers mainly dilute these waters. Si exhibits a low variability in response to the hydrological cycle because of the low contrast in concentration between the contributing tributaries (Fig. S1). A slight hysteresis for Ca and HCO_3 is observed upstream at the TAM, TAB and MAN stations (Solimões River), and a clear hysteresis is observed for SO_4 at the FAZ station (Madeira River). The variability in the Ca, HCO_3 and TDS concentrations remains low and does not explain the observed

hysteresis for these solutes at the OBI station. Indeed, considering a constant concentration of Ca, HCO_3 and TDS throughout the year (mean values in Table 3) at the upstream stations, we are able to reproduce the variability in concentration observed at the OBI station based on Eq. 2 (we considered a constant concentration and the monthly discharge at each station) with a maximum monthly error of $+9 \text{ mg L}^{-1}$ for TDS (Fig. 6), $+48 \text{ } \mu\text{mol L}^{-1}$ for Ca and $+88 \text{ } \mu\text{mol L}^{-1}$ for HCO_3 . The Si and K concentrations remain relatively constant at the upstream stations. The high hysteresis of SO_4 observed at the FAZ station explains a significant part of the SO_4 variability observed at the OBI station. Indeed, considering a constant SO_4 concentration upstream of the OBI station, the monthly SO_4 concentration is reproduced with a maximal error of $16 \text{ } \mu\text{mol L}^{-1}$ in July. Nevertheless, SO_4 contributes only to 4–9 % of the Amazon TDS over the course of the year. In

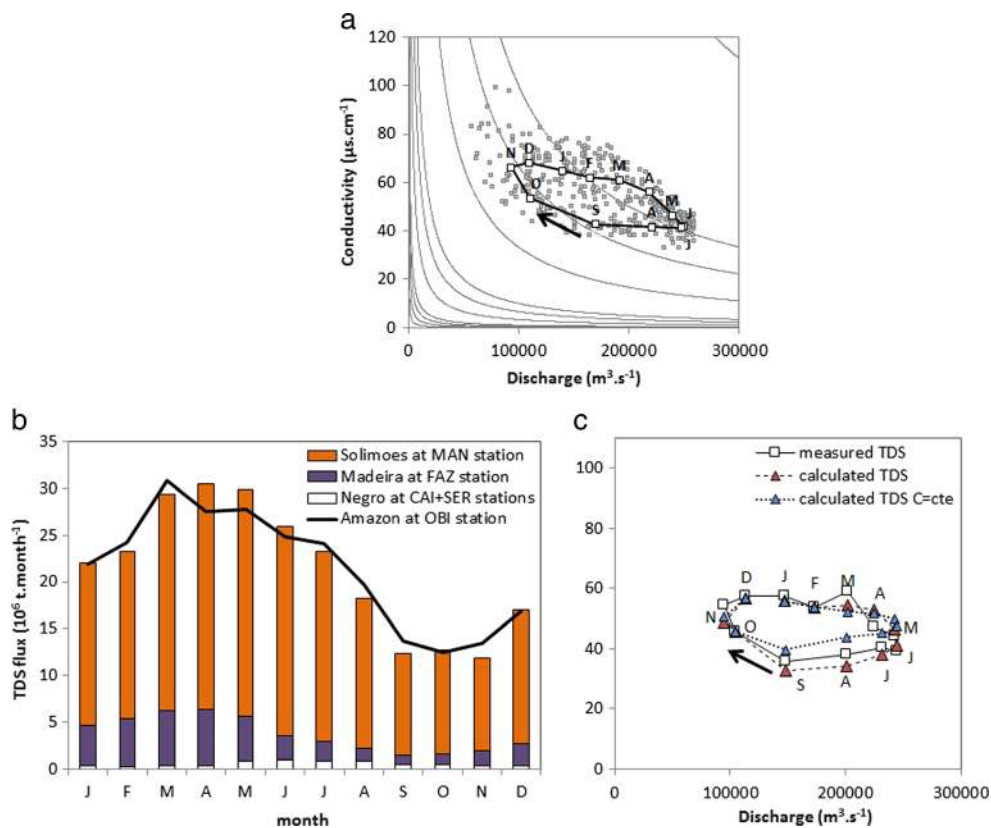


Fig. 6 **a** Ten-day frequency conductivity values (*small symbols*) and interannual monthly conductivity averages (*large symbols and lines*) at Óbidos as a function of discharge (September 1994 to December 2012 period). Simple dilution curves (concentration variability of a constant flux) are added for reference. **b** Comparison of the sum of the monthly TDS flux of the Solimões at the MAN station, Madeira at the FAZ station and Negro at the CAI and SER stations with the Amazon TDS flux at the

OBI station (January 2003 to December 2012 period). **c** Comparison of the mean monthly TDS concentration at the OBI station, the calculated Amazon TDS using fluxes (sum of the monthly TDS flux at the MAN, FAZ, SER and CAI stations divided by the monthly discharge at the OBI station) and the calculated Amazon TDS considering constant TDS concentrations at the MAN, FAZ, SER and CAI stations as a function of the mean monthly discharge (January 2003 to December 2012 period) at OBI

consequence, the dynamics of this solute do not contribute significantly to the observed TDS/conductivity hysteresis.

This result highlights the need for a monthly sampling frequency to accurately calculate the dissolved flux of the Amazon basin.

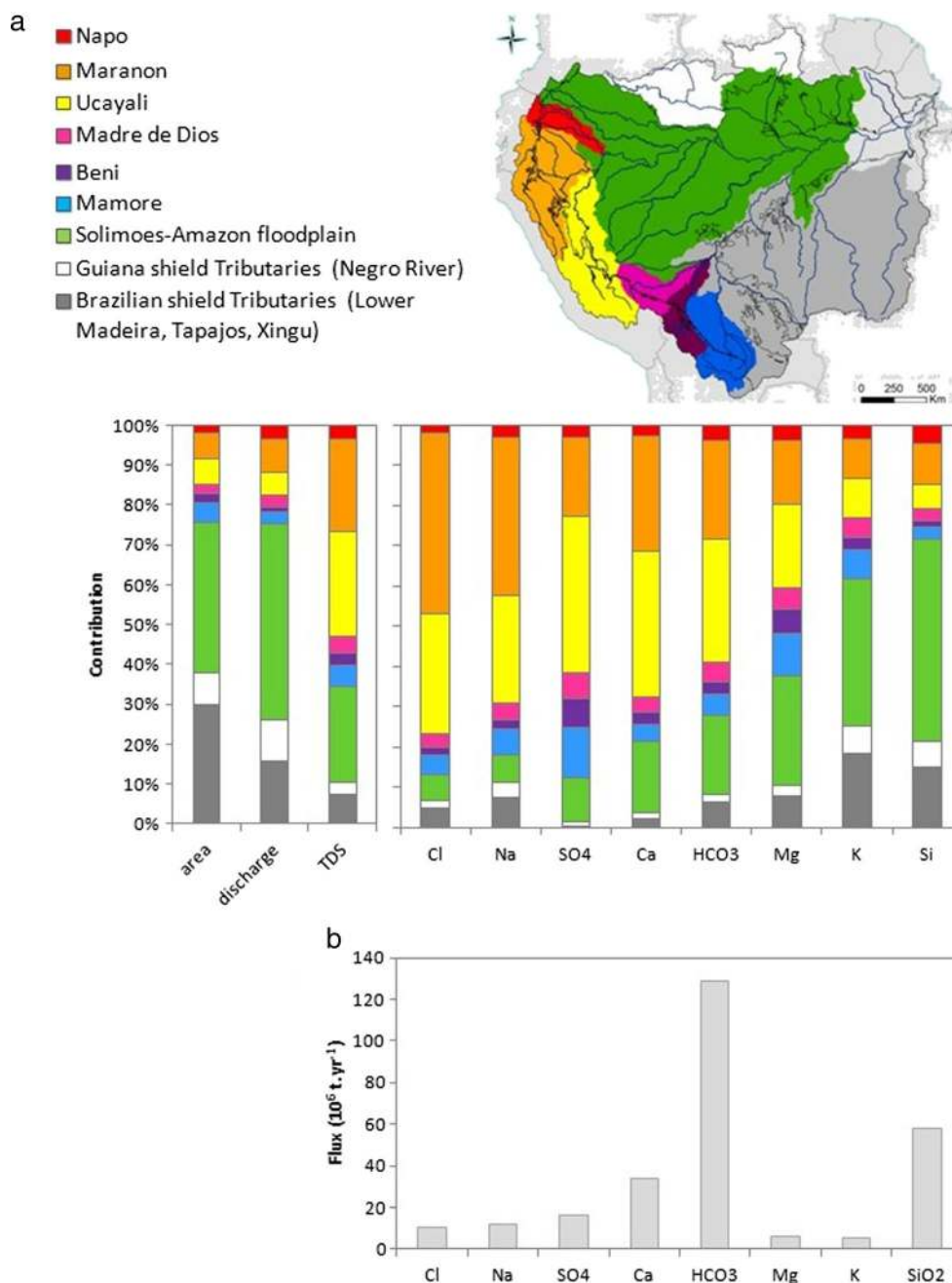
Annual flux calculation

The database allows us to estimate the solute fluxes from the different domains of the Amazon basin. The Amazon basin (Amazon flux at OBI + Tapajós Basin at ITA + Xingu basin at ALT) produced approximately $272 (\pm 54) \times 10^6$ t TDS year⁻¹ during the period 2003–2012. Using out four calculation methods (Table S1) based on the same database, results range between 263 and 278×10^6 t TDS year⁻¹ (Supplementary material S2).

The Andean tributaries (Napo, Marañon, Ucayali, Madre de Dios, Beni and Mamoré Rivers) contribute approximately 64 % of the Amazon TDS flux for only 26 % of the discharge and 25 % of the basin area (Fig. 7). The central plain (37 % of the basin area) and the shield tributaries (37 % of

the basin area) contribute approximately 25 and 11 % of the TDS flux, respectively (Table 4 and Fig. 7). The Marañon–Ucayali basins largely dominate the Andean TDS fluxes. These basins contribute approximately 50 % of the Amazon TDS flux for only 14 % of the Amazon discharge and 13 % of the Amazon basin area (Fig. 7). Nevertheless, this spatial flux distribution is not homogenous for all the elements (Table 4 and Fig. 7). The Marañon–Ucayali Rivers contribute 60–70 % of the Na, Ca and HCO₃ exports of the Amazon. Approximately 21 and 9 % of the Amazon HCO₃ flux are produced by the central plain area and the tributaries in the shields, respectively, whereas 70 % of this budget is produced by the Andean and foreland tributaries. Cl and SO₄ are mainly produced by the Andean and foreland sub-basins (83 and 88 % of the Amazon flux, respectively). Even if the Andean basins dominate the Mg flux, the contribution from the Solimões central plain and the shields is significant (37 % of the Amazon flux). In contrast with the other elements, the Amazon central plain and the shields are the main sources of K and Si (Table 4 and Fig. 7).

Fig. 7 **a** Contribution of each sub-basin of the Amazon basin to the surface area, the discharge and the dissolved load production. (TDS = Ca + Mg + Na + K + Cl + SO₄ + HCO₃ + SiO₂ in mass unit). **b** Total flux of each major element from the Amazon



Discussion

Relationships between concentration and discharge

Solutes delivered by evaporite dissolution

Cl, Na and SO₄ are mainly produced by the Andean and foreland Solimões sub-basins. In these contexts, the main sources of these elements are the dissolution of evaporites (Cl, Na and SO₄) (Stallard and Edmond 1983; Moquet et al. 2011), the anthropogenic inputs in the El Tigre basin before February 2009 (Cl) (Moquet et al. 2014b) and pyrite oxidation (SO₄)

(Moquet et al. 2011). The remaining Cl, Na and SO₄ produced by the central plain and the shields are mainly derived from atmospheric inputs (Cl, Na and SO₄), silicate weathering (Na) and, in the shields, from pyrite oxidation (SO₄) as evaporite are nearly absent their (Stallard and Edmond 1983). However, in the Solimões Andean and foreland basins, these elements can be considered as proxies for evaporite dissolution.

After February 2009 (when anthropogenic inputs stopped), *b* value is close to -1 for these elements controlled by evaporites inputs (Moquet et al. 2014b). This implies that the dissolved flux released by evaporite dissolution is relatively constant and insensitive to discharge variability, as also observed

Table 4 Dissolved load flux calculation by geomorphological setting

Sub-basins classified by geomorphological setting		Mean discharge $10^3 \text{ m}^3 \text{ s}^{-1}$	Area 10^6 km^2	Ca 10^6 t year^{-1}	Mg	Na	K	Cl	SO ₄	HCO ₃	SiO ₂	TDS 10^6 t year^{-1}
Andean basins	Flux	50	1.48	27	4	9	2	9	14	91	19	175
$F_{\text{TAB}}+F_{\text{PVE}}-F_{\text{VGR}}$	% Amazon	26 %	27 %	78 %	63 %	81 %	40 %	83 %	88 %	70 %	32 %	64 %
Shield tributaries	Flux	51	2.04	1.2	0.6	1.3	1.4	0.9	0.2	11	12	29
$F_{\text{SER}}+F_{\text{CAI}}+(F_{\text{FAZ}}-F_{\text{PVE}}+F_{\text{VGR}})+F_{\text{ITA}}+F_{\text{ALT}}$	% Amazon	27 %	37 %	4 %	10 %	12 %	25 %	9 %	2 %	9 %	20 %	11 %
Central plain tributaries	Flux	90	2.06	6.2	1.6	0.8	2.0	0.9	1.7	27	28	68
$F_{\text{OBI}}-F_{\text{FAZ}}-F_{\text{TAB}}-F_{\text{SER}}-F_{\text{CAI}}$	% Amazon	47 %	37 %	18 %	27 %	7 %	36 %	8 %	10 %	21 %	47 %	25 %
Amazon	Flux	191	5.58	34	6	12	6	11	16	129	58	272
$F_{\text{OBI}}+F_{\text{ITA}}+F_{\text{ALT}}$												

The formula used for the flux calculation is also reported

F_x annual flux of the considered major element at gauging station x

in the Southern Andes (Leon and Pedrozo 2015). A possible mechanism is that these inputs can be associated either with a constant groundwater flux after interaction with diapir domes or with constant flux from local resurgences of deep water. These deep saline waters would then be diluted by surface water during hydrological cycle.

Other sources

For Cl, SO₄ and Na in all other basins and for all elements in all contexts, the $C-Q$ relationship is shallow ($-0.1 > b > -0.5$) or nearly chemostatic ($0.1 > b > -0.1$). This implies that dilute rainfall water does not act as a simple diluter. Low dilution and chemostatic $C-Q$ relationships are observed in catchments of diverse sizes (10^1-10^6 km^2) (e.g. Kirchner 2003; Godsey et al. 2009; Clow and Mast 2010; Maher 2011; Basu et al. 2011; Guan et al. 2011; Moon et al. 2014; Stallard and Herndon et al. 2015). These relationships are the result of hydrological (mixing reservoirs, mixing tributaries) or chemical (kinetic or thermodynamic equilibrium) processes (Clow and Mast 2010; Maher 2011; Eiriksdottir et al. 2013) that influence the way solutes are produced and exported by watersheds. It has been suggested that an important control of the chemostatic or nearly chemostatic $C-Q$ response of a hydrosystem is the residence time of water (Maher 2010, 2011; Maher and Druhan 2014; Li et al., 2014; Maher and Chamberlain 2014). A chemostatic behaviour in rivers would be observed when the water has sufficient time to reach equilibrium with minerals. Various models have been able to reproduce the chemostatic behaviour observed in most settings (e.g. Godsey et al. 2009; Jawitz and Mitchell 2011). Based on catchments in the Colorado Rocky Mountains, Clow and Mast (2010) indicated that cation exchange and seasonal precipitation and dissolution of amorphous or poorly crystalline aluminosilicates are important processes that regulate the

concentration in a stream. At the scale of large rivers, the observed chemostatic behaviour can also be explained by the contribution of hydrological reservoirs and tributary inputs with different concentrations induced by different processes. The mix of these waters would result in a buffered signal with nearly constant concentration with increasing stream order (Creed et al. 2015).

For Ca, HCO₃ and TDS in the Andes, the chemostatic behaviour of the Marañon and Ucayali would therefore reflect more reactive mineralogy than that of the Napo and Madeira basins, where low dilution is observed (Figs. 4 and 5). This is consistent, at first order, with the fact that the carbonates, which are mainly restricted to the Marañon-Ucayali basins, are more reactive than silicates, which are the main sources of these elements in the Napo and the Upper Madeira (Moquet et al. 2011). Indeed, far from equilibrium, carbonates are highly reactive to weathering compared to silicate minerals (Lasaga and Berner 1998) and waters are able to rapidly reach equilibrium with respect to carbonate minerals.

In all Amazonian contexts, K and Si generally exhibit chemostatic behaviour. Godsey et al. (2009) observe the same phenomenon for Si in small catchments in the USA. They highlighted that the rates of reaction including these elements are very slow, implying that stream water should always be far from equilibrium with respect to the weathered material. The mechanism controlling chemostasis in this case may consequently be due to a ubiquitous source homogeneously distributed throughout the subsurface in the Amazon basin (clays or phytoliths for example) or to preferential contribution of the flushing of old water to the rivers.

Hysteresis behaviour at the OBI station

Previous studies on hysteresis of $C-Q$ relationships observed in rivers mainly focused on solid suspended matter

(e.g. Wood 1977; Walling and Webb 1982; Richey et al. 1986; Meade 1994; Jansson 2002; Guyot et al. 2005; Martinez et al. 2009; Filizola et al. 2011; Armijos et al. 2013), but this behaviour remains less documented for solutes.

Hysteresis for solutes $C-Q$ relationships is generally observed during storm events (e.g. House and Warwick 1998; Bowes et al. 2005) and is more rarely reported at the scale of large basins (Coynel et al. 2005; Ollivier et al. 2006). Hysteresis trends for dissolved solid concentrations are induced by time lags between dissolved solid and water contributions of reservoirs or tributaries (e.g. Ollivier et al. 2006). These time lags arise when different dissolution kinetics occur in the reservoirs and/or when the water transit times between reservoirs are different (e.g. House and Warwick 1998; Rose 2003). Solid suspended matter in rivers can also be a significant source or sink of some dissolved elements and can control the concentrations of those elements (Bowes et al. 2005). This last case is especially observed for bioreactive solutes, like phosphorus (Bowes et al. 2005; O'Connor et al. 2011), dissolved organic nitrates and dissolved organic carbon (Edokpa et al. 2015).

Although, in the Amazon, the hysteresis behaviour between TSS concentrations and discharge at the OBI station is well established (e.g. Richey et al. 1986; Meade 1994; Filizola and Guyot 2009; Filizola et al. 2011; Martinez et al. 2009), it is less well characterized for TDS concentration (Tardy et al. 2005; Bustillo et al. 2010) and never with a high frequency conductivity allowing a precise characterization of the phenomena.

Both TDS and TSS hysteresis phenomena are clockwise, but their relative variability in response to the hydrological cycle is different (Fig. 8a, b). Indeed, the relative amplitude of TSS (9.8 to 255 mg L^{-1}) is higher than that of conductivity (26 to 99 $\mu\text{S cm}^{-1}$), whereas the daily discharge (over the days of sampling) varies from $52,500$ to $259,600$ $\text{m}^3 \text{s}^{-1}$. In response to the hydrological cycle, during the decreasing and low water stages, high variability in conductivity is recorded when TSS is almost constant and minimal. During the increasing water stage, the TDS concentration remains high because of the high input of the Solimões River, whereas TSS increases. In the March–April–May (MAM) periods, during the second half of the increasing water stage, TDS and TSS decrease together because of the dilution effect of the Negro inputs for both TSS and TDS concentrations (Fig. 8c).

TSS and TDS concentrations respond to different mobilization processes (as generally observed in most rivers; e.g. Négrel et al. 2007; Sondag et al. 2010), and they are not produced in equivalent proportion by the Amazon tributaries. In the Amazon, the TSS dynamics has been explained by the combination of various processes such as the ‘depletion’ or ‘exhaustion’ effect (e.g. Meade 1994), the lag in time of tributary inputs (Filizola and Guyot 2009; Filizola et al. 2011) and

the backwater effect (Meade et al. 1991; Espinoza et al. 2013), but this behaviour remains debated.

In contrast with TSS, the hysteresis observed for TDS is explained by a simple mixing model. The three main tributaries (Solimões, Madeira and Rio Negro) show different average TDS concentrations with low fluctuation during the hydrological cycle. Consequently, hysteresis of TDS concentration at OBI arises mainly because of offsets between the contributions of the main tributaries during the hydrologic cycle (Fig. 6).

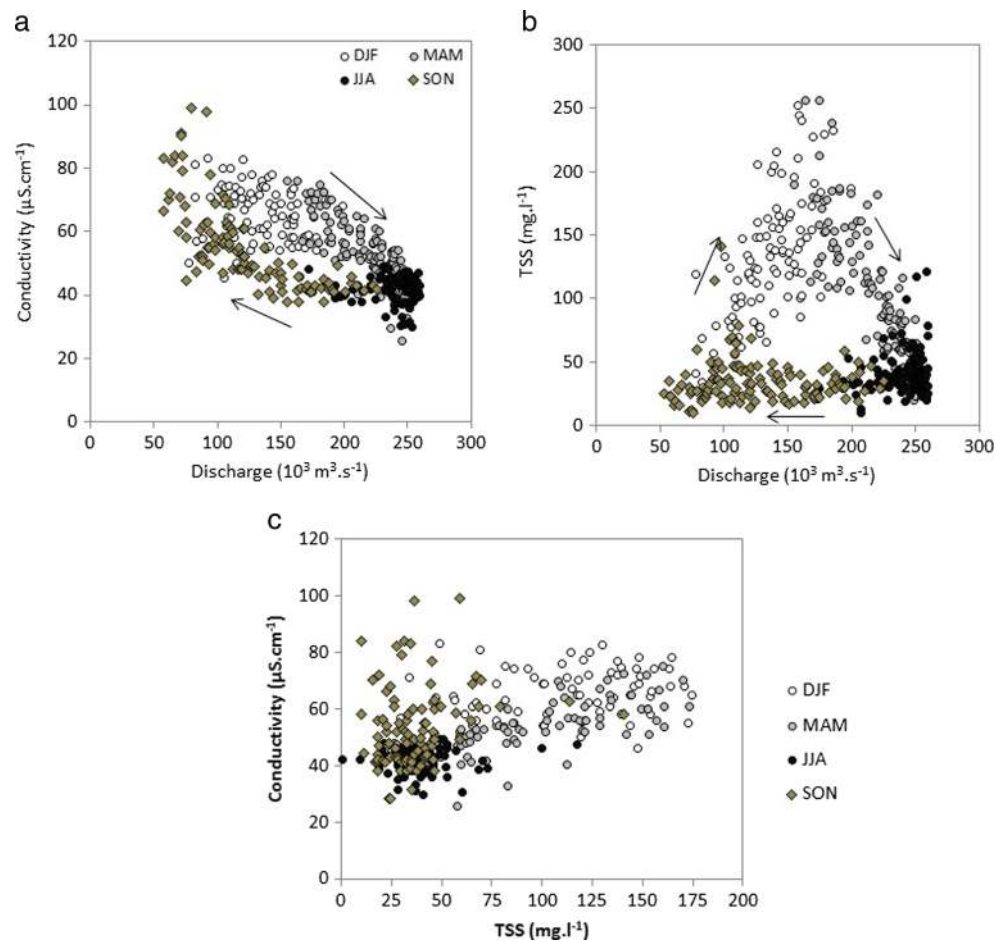
Origin of the dissolved load throughout the Amazon basin

The Andean inputs

The Andean (including foreland) basins produce approximately 172×10^6 t TDS year^{-1} and contribute approximately 64 % of the TDS of the Amazon basin (Figs. 7 and 9). Even though the Andean TDS contribution to the Amazon exports was identified early on as the main solute source of the Amazon River (Gibbs 1967a, b; Stallard and Edmond 1983), estimates of its relative contribution to TDS and solutes fluxes have rarely been reported (McClain and Naiman 2008; Mortatti and Probst 2003). Based on eight samples along the hydrological cycle on the main tributaries of the Solimões–Amazon River, Mortatti and Probst (2003) estimated that the Andean basins contributed up to 80 % (201 t TDS year^{-1}) of the Amazon TDS production. Our estimates are lower than these estimates. The difference can be attributed to both differences in sampling frequency and to the spatial distribution of the sampled stations. The high sampling frequency and the distribution of the SO-HYBAM framework stations allow for precise quantification of TDS and solute fluxes and of the relative contribution of the Andes and central plain.

The TDS specific flux from the Andes (118×10^3 t $\text{km}^{-2} \text{year}^{-1}$) is larger than that from the plain (33×10^3 t $\text{km}^{-2} \text{year}^{-1}$) and the shields (14×10^3 t $\text{km}^{-2} \text{year}^{-1}$) for similar runoff values ($1,070$, $1,375$ and 790 mm year^{-1} for the Andes, central plain and shields, respectively) (Fig. 9b). Specific fluxes are a function of concentration and of runoff. We previously showed that the temporal variability of most of the solute fluxes is mainly controlled by discharge variability. Nevertheless, as the mean runoff across these three geomorphological domains is similar, the Andes–sedimentary areas–shield TDS-specific flux (and flux) gradient observed throughout the basin is controlled by the contrast in TDS concentration values rather than by runoff (and discharge) variability. This observation shows that the total weathering flux of the Amazon is primarily controlled by the geomorphological/geological setting rather than by the spatial variability of the water supply (Figs. 7 and 9). This result confirms the high sensitivity of the Andean environment to weathering processes compared with sedimentary areas and

Fig. 8 Ten-day frequency conductivity values (**a**) and TSS concentration (**b**) as a function of discharge at the Óbidos station (September 1994 to December 2012 period). **c** Conductivity as a function of TSS concentration



shields. Indeed, in orogenic areas such as the Andes, high slopes and physical erosion favour the contact between minerals sensitive to weathering processes (carbonates, evaporites and primary silicate minerals) and water (Stallard 1985; Raymo and Ruddiman 1992; Dupré et al. 2003; Milliman and Farnsworth 2011). Furthermore, the TDS b value decreases from Andean basins to floodplain areas (Fig. 5). Torres et al. (2015) observed also this phenomenon for solutes derived from silicate weathering in the Andean and Andean foreland of the Madre de Dios basin (Madeira River). They explained this observation by the topography and/or erosional regime differences between Andes and Andean foreland contexts. In particular, in mountains, the high erosion would favour the abundance of reactive mineral and the fluid transit time in fractured bedrocks would be longer than in plain, while foreland floodplain is composed of thick soils depleted in reactive minerals and, possibly, shorter fluid transit time.

Role of the plains in the Amazon weathering budget

The central plain area is currently active in terms of the TDS, HCO_3 , Ca, Mg, K and Si production of the Amazon basin (Fig. 7). Because of the absence of evaporites and carbonates,

there silicate weathering and atmospheric inputs are the main sources of dissolved load in this area. The atmospheric inputs do not significantly influence the HCO_3 and Si of rivers (Berner and Berner 1987; Stallard and Edmond 1983), and with the exception of the Negro River for Mg (Gaillardet et al. 1997), they influence less than 10 % of the Ca, Mg and K production (Gaillardet et al. 1997; Stallard and Edmond 1983). These elements are consequently released to the river from silicates directly by weathering processes (Ca, Mg, HCO_3 , K and Si) or indirectly through plant uptake and decay (K and Si). This implies that the central plain area is currently active in terms of silicate weathering and has a significant influence on the production of dissolved load in the Amazon basin. This is well illustrated by the budget of HCO_3 . In the Andean and foreland area, 75 (61–83)% of the annual HCO_3 flux derives from carbonate weathering (Moquet et al. 2011). Because the sub-basins of the central plain and shields are mainly underlain by silicate lithology and because rainfall inputs are negligible for HCO_3 production in rivers, the HCO_3 produced in this area is mainly derived from CO_2 consumption associated with silicate weathering processes (CO_2 sil). Thus, the Andean sub-basins, the central plain sub-basins and the shield sub-basins are, respectively, responsible for approximately 41 %

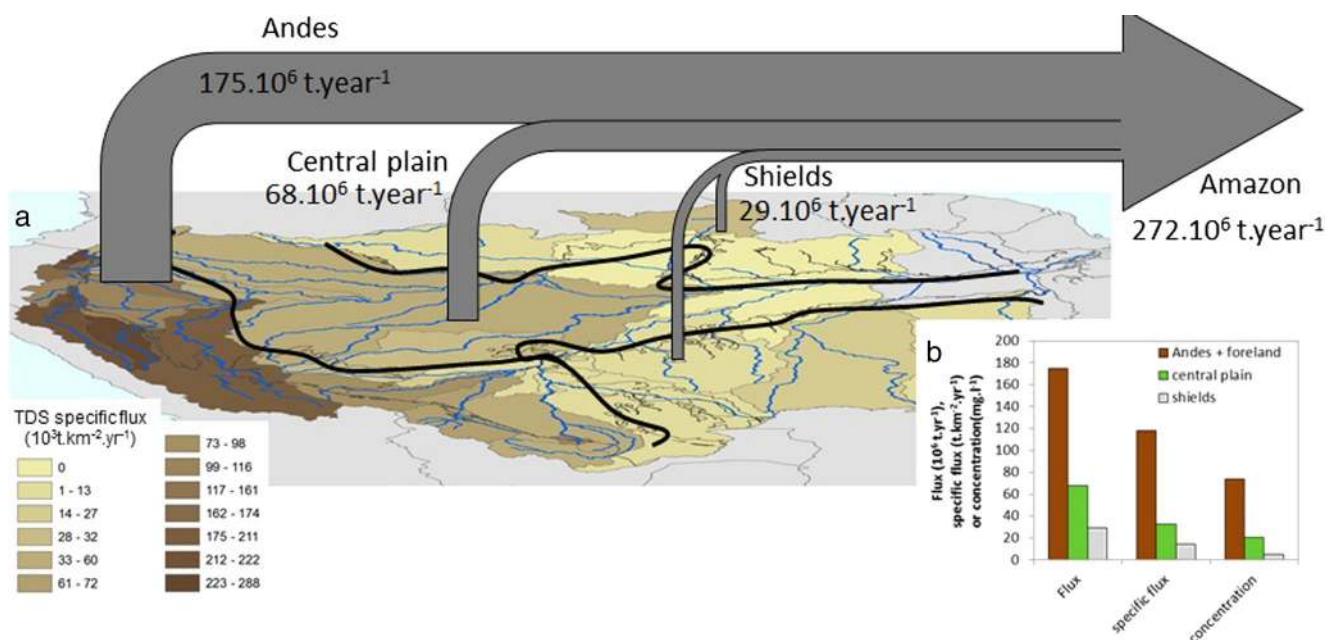


Fig. 9 **a** Specific flux of the dissolved load in the Amazon basin and TDS flux budget as a function of the main geomorphological contexts. **b** TDS flux, specific flux and mean concentration for the Andes, central plain and shields

(33–45), 41 % (33–45) and 18 % of the HCO_3^- flux derived from silicate weathering ($\text{HCO}_3^- \text{ sil}$) (or $\text{CO}_2 \text{ sil}$ consumption). Following this calculation, the relative contribution of Andean $\text{CO}_2 \text{ sil}$ consumption appears much lower than previous estimates (Mortatti and Probst 2003; Moquet et al. 2011). Nevertheless, this first-order estimate needs to be confirmed by further mass balance calculations allowing discrimination of the solute sources (atmospheric, evaporites, silicates, carbonates, anthropogenic inputs) (e.g. Gaillardet et al. 1997, 1999; Moon et al. 2014).

Our number shows that silicate weathering processes occurring in the sedimentary areas of an active mountain range can be significant at the scale of the basin, as shown by previous studies in the Amazon basin (Moquet et al. 2011; Bouchez et al. 2012, 2014) and the Brahmaputra basin (Galy and France-Lanord 1999; Lupker et al. 2012). The weathering budgets of these sedimentary area need to be taken into account for long-term carbon cycle modelling to estimate the long-term CO_2 variability and the links between orogenesis, climates and chemical–physical erosion processes (e.g. GEOCARB, Berner and Kothavala 2001; COPSE, Mills et al. 2014; WITCH, Godderis et al. 2006; CIDRE, Carretier et al. 2014).

The shields

The shields rivers exhibit both the lowest TDS concentrations and the lowest TDS-specific fluxes observed of the studied stations (Fig. 9b). This observation is consistent with the previous observations of Stallard and Edmond (1983) and Gaillardet et al. (1997). Indeed, the formation of thick tropical

soils would protect the bedrock from chemical weathering (Stallard 1985). This effect is particularly marked in the Upper Negro basin (SER station), where very low TDS concentrations are recorded ($\sim 6 \text{ mg L}^{-1}$) in comparison with the three other tributaries in the shields (the Branco, Tapajos and Xingu Rivers) (Table 3). The high runoff received by the upper Negro River compensates partially for the low concentration making TDS specific fluxes relatively high. Indeed, the TDS specific flux delivered by this area ($\sim 13 \times 10^3 \text{ t km}^{-2} \text{ year}^{-1}$) is similar with the TDS specific flux measured in the Tapajos ($19 \times 10^3 \text{ km}^{-2} \text{ year}^{-1}$) and Xingu Rivers ($13 \times 10^3 \text{ t km}^{-2} \text{ year}^{-1}$) but is low compared with that of the Branco River ($31 \times 10^3 \text{ km}^{-2} \text{ year}^{-1}$).

The Upper Negro (SER station) is characterized by podzols/arenosols (more than 40 % of the area; Dijkshoorn et al. 2005), which are mainly composed of quartz and have low sensitivity to weathering processes. In contrast, podzol/arenosol formations represent less than 3 % of the surface of the other shield basins. Even if the deep soils of the area are known to be poor in mobile elements (Stallard and Edmond 1983, 1987), these soils are currently submitted to weathering processes and are still able to produce dissolved mobile elements. Indeed, due to their surface area and their contribution of discharge to the Amazon, the shields contribute significantly to the TDS production (11 % of the Amazon flux), especially for Si and K production (20 and 25 % of the Amazon flux, respectively). This observation highlights that the main parts of the shield areas have a low impact on the total weathering budget of the Amazon basin, although they are chemically active environments.

Annual Amazon dissolved load export to the oceans

The Amazon basin (Amazon flux at OBI + Tapajós Basin at ITA + Xingu basin at ALT) produced approximately 272×10^6 t TDS year⁻¹ during the monitored period of 2003–2012 (Fig. 9). This value is close to previous estimates ranging from 251 to 292×10^6 t TDS year⁻¹ (Gibbs 1972; Berner and Berner 1987; Gaillardet et al. 1997, 1999; Mortatti and Probst 2003; Milliman and Farnsworth 2011; Sanchez et al. 2015). The flux for each element is also generally commensurate with the estimates of Gibbs (1972) and Mortatti and Probst (2003). The Amazon TDS solute production is dominated by HCO₃, Si and Ca, which represent 48, 21 and 12 % of the TDS, respectively (mass unit). Using the annually exported TSS ($600\text{--}800 \times 10^6$ t year⁻¹; Guyot et al. 2005; Martinez et al. 2009; Filizola et al. 2011), the TDS flux corresponds to 25–31 % of the total solid material carried by the Amazon River.

Using the Berner and Berner (1987) and Milliman and Farnsworth (2011) estimates of global continental TDS exportations ($3,843 \times 10^6$ and $3,800 \times 10^6$ t TDS year⁻¹, respectively), the Amazon River contributes approximately 7 % of the total dissolved load exported to the oceans. As highlighted by these authors, the Amazon River ranks as the most important basin in terms of dissolved load flux before the Changjiang (180 t TDS year⁻¹; Milliman and Farnsworth 2011) and has a dissolved yield (49 t TDS km⁻² year⁻¹) similar with the world average (~ 40 t TDS km⁻² year⁻¹; Milliman and Farnsworth 2011). Individually, the Solimões, the Madeira and the rivers of the shields (Negro + Tapajós + Xingu) contribute up to 5.6 % (214 t TDS year⁻¹), 1.0 % (39 t TDS year⁻¹) and 0.6 % (22 t TDS year⁻¹) of the global TDS flux, respectively. The Solimões itself is the first world basin in terms of the flux of river dissolved load. Following the hierarchy of 62 largest rivers of Gaillardet et al. (1999), the TDS flux of the Madeira River is similar with that of the Rhine (18th position), and the TDS fluxes delivered by the tributaries from the shields are together commensurate with those of the Don and Nelson River TDS fluxes (26th and 27th positions, respectively). The Solimões exhibits higher TDS specific flux values (97 t TDS km⁻² year⁻¹) than the global average and is similar with that of the Ganges Brahmaputra basin (which ranges as the third highest dissolved load of world rivers; Milliman and Farnsworth 2011), whereas the Madeira and the basins of the shields exhibit lower values than the world average (29 and 17 t TDS km⁻² year⁻¹, respectively). The Ca, Mg, Na, Cl, SO₄ and HCO₃ carried by the Amazon River to the oceans correspond to 5–8 % of the world riverine fluxes, and the dissolved Si and K exports correspond to 10–12 %.

In the 2003–2012 period, only the Cl concentration appears to show a slight decrease between the period before and after February 2009 due to the decrease in contamination by oil extraction activities identified throughout the northern foreland (Tigre River at NYO) (Moquet et al. 2014b). The other

elements do not show a significant interannual trend during the 10 years of monitoring at the Óbidos station.

Potential effects of further climate changes

During the last decades, the Amazon basin has been affected by climate changes in term of both the annual distribution of rainfall and increasing frequency of extreme events (Espinoza et al. 2009b; Lavado Casimiro et al. 2012). From global and regional climate models, climate projections predict that these trends will intensify in the future (IPCC report: Magrin et al. 2014). Projections reported by the IPCC (Magrin et al. 2014) agree that the basin will experience a significant increase in temperature. The projections for rainfall are affected by large uncertainties, but some general trends can be extracted on a regional scale. The northwestern part of the basin would experience higher precipitation, whereas central, southern and eastern Amazonia would be subjected to drier conditions (Giorgi and Diffenbaugh 2008; Bombardi and Carvalho 2009; Marengo et al. 2011; Magrin et al. 2014). Increases in the frequency and intensity of extreme events would accompany these general trends. Following these scenarios, if the observed $C\text{--}Q$ relationships at the scale of the hydrological year are valid for decadal evolution of discharge, the Amazon TDS fluxes would increase in the future because the main source of TDS is located in the northwestern part of the basin. In particular, the dissolved fluxes of the Amazon mainly associated with carbonate weathering in the Marañon and Ucayali basins would increase proportionally to the discharge, whereas the Madeira TDS fluxes would decrease. Cl and Na fluxes, controlled by evaporite inputs, would not be affected by projected climate changes. Si and K fluxes would vary proportionally with the mean rainfall variation of the Amazon basin. As a perspective and as performed by Von Blanckenburg et al. (2015) for the Last Glacial Maximum based on Maher and Chamberlain (2014) $C\text{--}Q$ model, further combination of derived $C\text{--}Q$ relationships with climate model output projections of Amazon river discharge (and its tributaries) would allow to assess future changes in flux due to the anthropogenically altered climate state.

Nevertheless, these predictions assume that the $C\text{--}Q$ relationships are stationary through time. However, weathering rates depend not only on water availability but also on a number of other parameters such as lithology, temperature, erosion rates, abundance of organic acids and vegetation (Goudie and Viles 2012 and references within). Moreover, it is likely that the $C\text{--}Q$ relationships determined for a given range of Q are not stable for extreme conditions (Maher 2010). Further studies on small basins and on basins undergoing extreme climatic conditions need to be performed to better constrain these laws, to model chemical weathering processes and to better predict the future evolution of dissolved matter exportation from the Amazon basin.

Conclusion

In the period 2003–2012, the Amazon basin exported an average of 272×10^6 t year⁻¹ of dissolved load, which corresponds to nearly 7 % of the TDS flux annually exported from the continents to the oceans. The Amazon TDS exports are mainly composed of HCO₃, Si and Ca. Ca and HCO₃ are mainly produced by carbonate weathering in the Andean basins and especially in the Marañon and Ucayali basins (representing approximately 50 % of their fluxes), whereas Si production is mainly controlled by the spatial distribution of rainfall throughout the basin. A TDS concentration gradient is observed according to the following geomorphologic hierarchy: Andean basins > sedimentary areas > non-podzolic shields > podzolic shields. Due to its surface and its relative contribution to the Amazon, the water flux from the central plain significantly contributes to the dissolved production of the basin, with an export of approximately 25 % of the total TDS flux. This result confirms that sedimentary areas are currently undergoing weathering processes. Further analysis needs to be performed to accurately discriminate the TDS sources.

Over the main Amazon tributaries, the variability of TDS fluxes is mainly controlled by discharge variability in response to the hydrological cycle. Evaporite dissolution produces a nearly constant flux throughout the year, whereas the TDS fluxes released by other sources (carbonate weathering, silicate weathering, biosphere and atmospheric inputs) are mainly controlled by variability in discharge. For the first time, the hysteresis behaviour observed for TDS (conductivity) at the Óbidos station is described. This phenomenon is mainly explained by the time lag in discharge seasonality of the three main tributaries (Solimões, Madeira and Negro), which have contrasted TDS concentrations. If the *C–Q* relationships determined throughout the basin remain stable, further changes in rainfall distribution and intensity would consequently have a direct impact on the variability of TDS flux. In particular, an increase in rainfall over the western and northwestern sub-basins underlain by carbonates would increase the exportation of Ca and HCO₃, whereas the dissolved Si exportation would be nearly proportionally affected by the changes occurring over the Amazon basin surface. Nevertheless, the main processes controlling the *C–Q* relationships of solutes throughout the basin are not identified. Further studies are still required to identify the main physico-chemical processes controlling these empirical laws, to better constrain their limitations and to quantify the effect of extreme events on these solute export budgets.

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