



Scholars Research Library

Archives of Applied Science Research, 2016, 8 (11):1-6  
(<http://scholarsresearchlibrary.com/archive.html>)



Scholars Research Library  
ISSN 0975-508X  
CODEN (USA) AASRC9

## Analysis of Polycyclic Aromatic Hydrocarbons (PAHs) in road runoff water from high traffic area of Guwahati City, Assam, India

Nilutpala Dutta<sup>1\*</sup> and Yamin Hassan<sup>2</sup>

<sup>1\*</sup>Research Scholar, Assam Down Town University, Assam down town University, Guwahati, India

<sup>2</sup>Assistant Professor, Chemistry Assam Down Town University, Assam down town University, Guwahati, India

### ABSTRACT

Road runoff water samples were collected for the analysis of eight PAHs during first flush and aftermath in monsoon season of the year 2014-15 from 12 sites associated with high traffic activities around the clock. All the sites were selected for investigation to represent the high traffic activity areas of Guwahati city, Assam, India. Eight common environmental PAHs [Phenanthrene-(PHE), Benzo(K)FLUORANTHENE(-BKF), Benz(a) Pyrene-(BAP), Benzo(ghi)Pyrene-(BGHIP), Benzo(a) Anthracene/Chrysene-(BAA/CHR), Fluoranthene-(FLT), Pyrene-(PYR), Anthracene-(ANT)] were found to be distributed and the concentration of PAHs was determined by using HPLC technique. Polycyclic aromatic hydrocarbons (PAHs) are potentially mutagenic and carcinogenic substances occurring at various concentrations in atmosphere, soils, waters, and sediments. PAHs, inherited both from natural and anthropogenic processes, are persistent organic pollutants (POP) due to their chemical stability and biodegradation resistance.

The present data indicates that the total average of PAHs over the investigated sites was ranged from 0.005 ng/l to 0.057 ng/l. On individual scale, the highest concentration was 0.057 ng/l and 0.053 ng/l for BKF and ANT. The investigation showed the carcinogenic content of PAHs(BKF) was found in all the investigated runoff water samples. The increase of road transportation, and of industrial and activities has led to a notable buildup of PAH amounts in the environmental media.

**Keywords:** Pyrene, Benz(a) Pyrene, High traffic activity area, Polycyclic aromatic hydrocarbons (PAHs), high performance liquid chromatography (HPLC)

### INTRODUCTION

A good quantity of contaminates in rain water runoff has become an increasing concern in rain water utilization. Storm water runoff from urban areas is a significant source of pollution to inland water bodies such as streams, rivers and lakes [1]. Rain water utilization plays a very important role as one of the best counter measures against water shortage in urban areas. It can overcome a shortage of water supply and in the meanwhile is very effective for runoff control [2-4]. Non potent pollution resulting from urban surface runoff was recognized as one of the major cause of quality deterioration in receiving water bodies [5-7]. Storm water runoff not only flows into the river but also rubbish, animal droppings, chemicals, fertilizers, oils, soil and anything that is placed in or washes into street gutters can end up in the river and polluting the environment [1]. The major sources of PAH in stormwater are the wear and leaching of asphalt, tire wear, drips of crankcase oil, and vehicular exhaust [8-10]. The most abundant PAHs in storm water are phenanthrene, anthracene, fluoranthene and pyrene [11]. Gonzalez et al. (2000) reported that these four PAHs accounted for 85% from a total 16 PAH in stormwater. Several PAHs are carcinogenic and mutagenic [12], and are included in the United States Environmental Protection Agency (USEPA) and EU priority pollutants list. According to a Danish study, runoff-contaminated sediments from a stream and a retention basin were polluted with heavy metals and PAH, and the sediment and the pore water were toxic to algae [13]. Boxall and Maltby (1995) and Maltby et al. (1995) found that organic compounds were the source of the toxicity in runoff-contaminated sediment and that the fraction of three to five-ring PAHs were the major toxicants [14,15]

### MATERIALS AND METHODOLOGY- ANALYSIS OF PAHS IN ROAD RUNOFF WATER

#### Study Area

Guwahati is the principal city of the entire Northeastern India and was chosen as the study area for the present work.

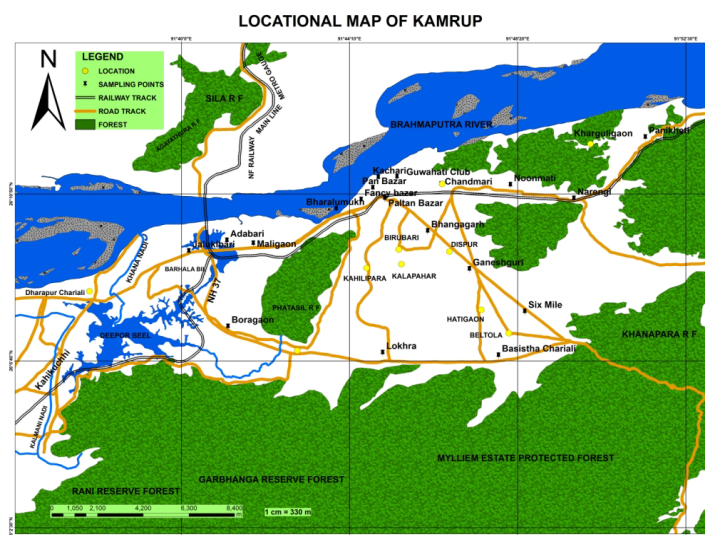
Guwahati is located approximately along 26° 11' N latitude and 91° 45' E longitude. It is 54.75 m above the mean sea level, covering about 24 km in the East-West direction and about 9 km in the North-South direction. The mighty Brahmaputra flows along the Northern boundary of the city while the Southern and Eastern boundaries are made by several hill ranges, which are extensions of the Khasi Hills. The Jalukbari-Azara plain makes the Western boundary of the city. The expert plan of the city also covers Amingaon and North Guwahati on the Northern side of the Brahmaputra. Structurally, this region is situated on the 50 m thick alluvium of the middle Brahmaputra valley. The city is situated on an outcrop of the stable rocky foundation of the Shillong Plateau and the floodplains of Brahmaputra confronting each other. Landforms within the city are therefore unique with dissected hills (originally part of the Shillong Plateau), plain areas and natural lakes (the beels), swamps and the mighty river Brahmaputra.

**Collection of samples**

The water samples have been collected in pre-cleaned 2 L polyvinyl containers from 12 different locations of Guwahati based on high traffic density areas during first flush and aftermath of first flush in the monsoon season of 2014-15. Necessary care has been taken to prevent contamination of the samples during transportation to the laboratory, storage, and analysis [16,17]. The geographical locations of the sampling points are given in Table 1. The base map of the study area is shown in Figure 1.

**Table 1.** The geographical coordinates of road runoff water sampling points during 2014-15 monsoon season

Sl No	Sampling Code	Geographical Coordinates
1	RW 1	N 26°11'00.9" and E 091°46'26.9"
2	RW 2	N 26°08'57.1" and E 091°47'08.5"
3	RW 3	N 26°07'14.1" and E 091°49'18.5"
4	RW 4	N 26°05'56.7" and E 091°52'35.4"
5	RW 5	N 26°09'53.8" and E 091°46'06.5"
6	RW 6	N 26°10'43.3" and E 091°45'02.8"
7	RW 7	N 226°09'32.9" and E 091°41'47.5"
8	RW 8	N 26°09'29.9" and E 091°40'52.5"
9	RW 9	N 26°11'05.9" and E 091°45'25.4"
10	RW 10	N 26°06'43.8" and E 091°47'52.6"
11	RW 11	N 226°09'26.5" and E 091°40'25.0"
12	RW 12	N 26°02'42.2" and E 091°52'04.9"



**Figure 1.** Base map of the study area.

**MATERIALS AND REAGENTS FOR PAH DETERMINATION IN ROAD RUNOFF WATER**

The water samples were stored at +4°C with the addition of 0.05% (w/v) sodium azide (NaN<sub>3</sub>). For preconcentration of PAHs by solid phase extraction (SPE) Bakerbond C18 microcolumns (6.0 mL, 1000 mg) and Speedisk Bakerbond SPE-50 (C18) from J.T.Baker were used. Additionally single-use microfilters type Millex-HV (0.45 μm) from Millipore were used. For conditioning of SPE columns and speedisks, removal of PAHs and separation by liquid

chromatography solvents for HPLC separation (HPLC grade: water, acetonitrile, dichloromethane, methanol, ethanol, and isopropanol) from J.T. Baker were used. As a standard the mixture of 16 PAHs (naphthalene, acenaphthene, acenaphthylene, fluorene, phenanthrene, anthracene, fluoranthene, pyrene, benzo[a]anthracene, chrysene, benzo[b]fluoranthene, benzo[k]fluoranthene, benzo[a]pyrene, dibenzo-[a,h]anthracene, benzo[g,h,i]-perylene, indeno[1,2,3-c,d]pyrene (purity 97.4-99.9%) in concentrations 10.0 mg/L of each from Ehrenstorfer (Germany) and standard of benzo[a]pyrene in concentration 100.0 mg/L ACN (purity 97.5%) from Sigma (USA) were used in experiments. All processes were done using single-use equipment and automatic pipets.

#### Preconcentration of PAHs by SPE method

The Bakerbond C18 microcolumns and Speedisks SPE-50 (C18) at the beginning were eluted (activated) by: 10.0 mL dichloromethane, 10.0 mL of isopropanol, 10.0 mL of ethanol, 10.0 mL of methanol and finally 10.0 mL of distilled and deionized water from Millipore system. The samples of water (1.0 L) were preconcentrated on Bakerbond C18 microcolumns (drinking water) or on Speedisk SPE-50 (C18) (lake or river water) with a flow rate of 2-3 mL/min. For preconcentration the SPE technique (SPE-12G from J.T. Baker) was used. The microcolumns and speedisk at the beginning were eluted by: 10.0 mL of distilled water, 10.0 mL of methanol, 10.0 mL of ethanol and 10.0 mL of isopropanol. Later were eluted by: 10.0 mL of ACN and 10.0 mL of CH<sub>2</sub>Cl<sub>2</sub> at a flow rate 2-3 mL/min. Both fractions eluted by ACN and CH<sub>2</sub>Cl<sub>2</sub> were collected and dried at inert gas flow at 28°C. The dried samples were resolved in 1.0 mL of CH<sub>2</sub>Cl<sub>2</sub>. For the measure of recovery of PAHs by SPE method 1.0 mL of solutions of 16 PAHs was added to 100 mL of CH<sub>2</sub>Cl<sub>2</sub>. For the measure of recovery of PAHs by SPE method 1.0 mL of solutions of 16 PAHs was added to 100 mL of distilled water. The samples were separated by preconcentrated by microcolumns Bakerbond C-18 and by Bakerbond speedisk SPE-50(C-18) as environmental samples and were determined by RP-HPLC method.

#### Analytical conditions for reversed-phase HPLC

Measurements were carried out using a HPLC system (Shimadzu LC-10 AD) with UV-visible detector. The analytical column was of 250 mm length and 4.6 mm i.d., packed with porous spherical RP-18 material (particle size 5µm). A guard column (10 mm long and 4.6 mm i.d.) preceded the analytical column. Acetonitrile-water mixture (70:30) was used as mobile phase at a flow rate of 1.0 ml per minute. Samples of 100 µl were injected into the column through the sample loop. A UV detector set at 254 nm for absorption was used for detection of the compounds. The data was processed with a CR 7A chromatopac data processor. Several dilutions corresponding to 0.2 to 20 ng absolute of synthetic standard mixture of individual components of phenanthrene, anthracene, fluoranthene, pyrene, benzo(a)anthracene, benzo(k)fluoranthene, and benzo(a)pyrene (supplied by Polyscience, USA) dissolved in HPLC grade acetonitrile was used for determining the retention data and for studying the linearity of the detector. The response was linear for a wide range of concentrations mentioned above. Using this method, PAH concentration of less than a nanogram in the sample could be quantified.

### RESULT AND DISCUSSION

In the present study eight PAHs were analysed namely Phenanthrene-(PHE), Benzo(K) Fluoranthene-(BKF), Benz(a) Pyrene-(BAP), Benzo(ghi)Pyrene-(BGHIP), Benzo(a) Anthracene/Chrysene-(BAA/CHR) Fluoranthene-(FLT), Pyrene-(PYR), Anthracene-(ANT). Among all these PAHs BAP, BAA/CHR, and BKF are carcinogenic according to CCME (Canadian Council of Ministers of The Environment). The total concentration of eight PAHs of first flush and aftermath runoff water is presented in Table 2 and Table 3. Among the eight PAHs only five PAHs namely PHE, BKF, FLT, PYR and ANT were detected. The concentration of BAP, BGHIP and BAA/CHR are found below detectable level. During first flush the concentration of detected PAHs were found in the range 0.009 ng/l to 0.057 ng/l (Table 2) while for the runoff samples collected aftermath of first flush was found in the range of 0.007 ng/l to 0.043 ng/l (Table 3). The mean concentration of five PAHs are given in Table 4. The mean concentration of all the detected five PAHs in the samples of first flush are more than the runoff water samples collected after first flush in the monsoon season as shown in Figure 2. The highest concentration were recorded for BKF (0.057 ng/l) and PHE (0.043 ng/l) as found in the Table 2 and Table 3. For all the five PAHs, the maximum concentration was recorded for the site RW12. The site RW12 witnessed many more coke processing units, brick industries, alloy manufacturing units, cement manufacturing units in the two sides of the road may also responsible for the high concentration of PAHs in the samples of sampling point RW12. According to Mastral AM et al. [18], Lee CW et al. [19], Khanfekr A et al. [20], Arulazhagan P et al. [21] and Fagbote EO et al. [22], other sources for the presence of PAHs in the environment include lubricating oils, atmospheric depositions, plants, domestic heating systems, petrol and diesel engines and various industrial activities.

**Table 2.** The total concentration of PAHs (ng/l) of first flush runoffwater during theyear 2014-15.

SI No	Sampling Code	PHE	BKF	BAP	BGHIP	BAA/CHR	FLT	PYR	ANT
1	RW 1	0.026	0.021	BDL	BDL	BDL	0.019	0.009	0.019
2	RW 2	0.017	0.012	BDL	BDL	BDL	0.014	0.026	0.023
3	RW 3	0.027	0.019	BDL	BDL	BDL	0.032	0.031	0.032
4	RW 4	0.037	0.043	BDL	BDL	BDL	0.021	0.021	0.036
5	RW 5	0.012	0.011	BDL	BDL	BDL	0.012	0.018	0.036
6	RW 6	0.047	0.014	BDL	BDL	BDL	0.031	0.036	0.042
7	RW 7	0.033	0.015	BDL	BDL	BDL	0.010	0.019	0.014
8	RW 8	0.034	0.022	BDL	BDL	BDL	0.029	0.036	0.035
9	RW 9	0.023	0.031	BDL	BDL	BDL	0.012	0.015	0.019
10	RW 10	0.032	0.033	BDL	BDL	BDL	0.020	0.039	0.030
11	RW 11	0.020	0.022	BDL	BDL	BDL	0.010	0.021	0.010
12	RW 12	0.052	0.057	BDL	BDL	BDL	0.034	0.044	0.053
	MIN	0.012	0.011	0.000	0.000	0.000	0.010	0.009	0.010
	MAX	0.052	0.057	0.000	0.000	0.000	0.034	0.044	0.053
	MEAN	0.030	0.025	0.000	0.000	0.000	0.020	0.026	0.029
	KURT	-0.171	1.360	0.000	0.000	0.000	-1.573	-1.081	-0.338
	SKEW	0.440	1.308	0.000	0.000	0.000	0.348	0.158	0.229

**Table 3.** The total concentration of PAHs(ng/l) of runoff water after first flash during the year 2014-15.

SI No	Sampling Code	PHE	BKF	BAP	BGHIP	BAA/CHR	FLT	PYR	ANT
1	RW 1	0.015	0.016	BDL	BDL	BDL	0.010	0.005	0.016
2	RW 2	0.007	0.008	BDL	BDL	BDL	0.008	0.019	0.021
3	RW 3	0.021	0.015	BDL	BDL	BDL	0.023	0.014	0.026
4	RW 4	0.032	0.016	BDL	BDL	BDL	0.012	0.007	0.021
5	RW 5	0.007	0.008	BDL	BDL	BDL	0.009	0.016	0.022
6	RW 6	0.039	0.005	BDL	BDL	BDL	0.023	0.017	0.033
7	RW 7	0.027	0.012	BDL	BDL	BDL	0.005	0.008	0.007
8	RW 8	0.024	0.022	BDL	BDL	BDL	0.019	0.017	0.031
9	RW 9	0.017	0.007	BDL	BDL	BDL	0.007	0.010	0.012
10	RW 10	0.024	0.024	BDL	BDL	BDL	0.014	0.009	0.024
11	RW 11	0.016	0.014	BDL	BDL	BDL	0.005	0.012	0.006
12	RW 12	0.043	0.025	BDL	BDL	BDL	0.027	0.032	0.040
	MIN	0.007	0.005	0.000	0.000	0.000	0.005	0.005	0.006
	MAX	0.043	0.025	0.000	0.000	0.000	0.027	0.032	0.040
	MEAN	0.023	0.014	0.000	0.000	0.000	0.014	0.014	0.022
	KURT	-0.462	-1.053	0.000	0.000	0.000	-1.094	2.732	-0.454
	SKEW	0.375	0.303	0.000	0.000	0.000	0.616	1.369	0.073

**Table 4.** The mean concentration of five PAHs of first flash and aftermath runoff water samples.

Sample	PHE(ng/l)	BKF(ng/l)	FLT(ng/l)	PYR(ng/l)	ANT(ng/l)
First Flush Runoff Water Sample	0.030	0.025	0.020	0.026	0.029
Runoff Water Sample After First Flush	0.023	0.014	0.014	0.014	0.021

**Statistical analysis**

Normal statistics of runoff water quality parameters of road runoff water samples are presented in Tables 2 and 3. Positive value of kurtosis was calculated for PYR for first Flush aftermath run off while for first flush run off samples it was found for BKF in 2014-15 season which show sharp distribution around the study area. Negative value of kurtosis for the remaining four PAHs

show a flat distribution around the study area. Positive value of skewness gives a distribution with a significant long right tail and negative value of skewness gives a distribution with a significant long left tail. Negative value of skewness was shown by BKF of runoff water samples collected after first flushwhereas other four PAHs showed the positive values but all the five PAHs of first flush run off samples have shown positive skewness.

### CONCLUSION

The most abundant PAHs in the runoff water samples were PHE, BKF, FLT, PYR and ANT. Except sampling point RW12 all other sampling sites are associated only with heavy traffic activities, so we may assume that mainly high traffic activities are responsible for the presence of detected PAHs in the study area. There was a significant difference between the mean concentrations of the first flush and aftermath of first flush runoff water samples as observed from Figure 2. This may be due to binding of the PAHs with runoff water supported by high affinity of PAHs for organic carbon, and most PAH can therefore be found attached to particles that have settled or are suspended in the water column [12,23]. Smith et al. (2000) reported in his study that the total concentration of PAHs was highest at the beginning of the runoff, the so called first flush, and decreases with time. The identification of the sources has shown that industrial sources in association with traffic sources may contribute a significant number of PAHs in the environment.

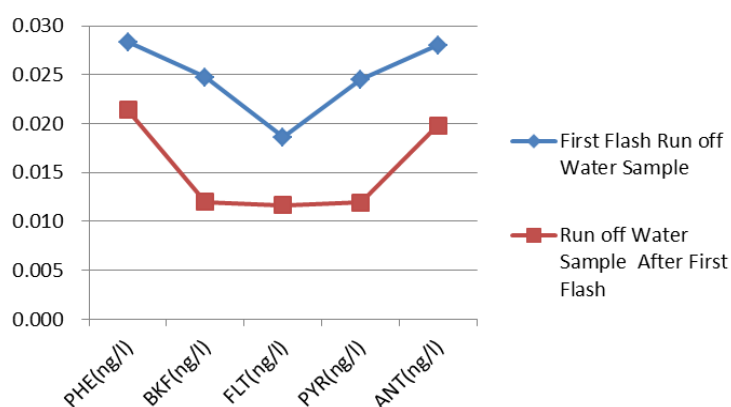


Figure 2. The comparison of mean concentration of five PAHs of runoff water samples of first flush and aftermath

### ACKNOWLEDGEMENT

The Authors acknowledge Assam Down Town University, GHY-26 for providing the permission and the facilities for carrying out the research work.

### REFERENCES

- [1] R A Allison, T A Walker, F H Schiew, I C O'Neill, T A McMahon, *CRC for Catchment Hydrology*, **1998**, 98, 6.
- [2] J Forster, *Water Sci. Technol*, **1996**, 33 (6), 39–48.
- [3] W He, I O Wallinder, C Leygraf, *Corros Sci*, **2001**, 43, 127–146.
- [4] J Smith, M Sievers, S Huang, S L Yu, *Water Science and Technology*, **2000**, 42, 383–388.
- [5] M Chang, C M Crowley, *Water Res Bull*, **1993**, 29(5), 777–783.
- [6] T Gnecco, C Berretta, L G Lanza, P L Barbera, *Atmospheric Research*, **2005**, 77, 60–73.
- [7] M C Gromaire-Mertz, S Garnaud, A Gonzalez, G Chebbo, *Water Sci Technol*, **1999**, 39 (2), 1–8.
- [8] H C A Brandt, P C De Groot, *Water Research*, **2001**, 35, 4200–4207.
- [9] J Brown, B Peake, *Science of the Total Environment*, **2006**, 359, 145–155.
- [10] J Neff, JMLA, *Polycyclic aromatic hydrocarbons in the aquatic environment: sources, fates, and biological effects* London, Applied Science Publishers Ltd, **1979**, 18, 92.
- [11] A Gonzalez, R Moilleron, G Chebbo, D Thévenot, *Polycyclic Aromatic Compounds*, **2000**, 20, 1–19.
- [12] ATSDR, *Toxicology profile for polycyclic aromatic hydrocarbons*, US Department of Health and Human Services, Public Health Service. Atlanta, GA, **1995**, 1, 6.
- [13] A M Christensen, F Nakajima, A Baun, *Environmental Pollution*, **2006**, 144, 621–625.
- [14] A B A Boxall, L Maltby, *Water Research*, **1995**, 29, 2043–2050.
- [15] L Maltby, A B A. Boxall, D M Forrow, P Calow, C Betton, *Environmental Toxicology and Chemistry*, **1995**, 14, 1093–1101.

- [16] APHA, *Standard Methods for Examination of Water and Waste Water*, American Public Health Association and American Water Environmental federation, USA, 1998, 20, 19.
- [17] R K Trivedy, P K Goel, *Chemical and biological methods for water pollution studies*, Environment publication, India, 1986, 12, 27-38.
- [18] A M Mastral, M S Callen, R Murillo, *Fuel*, **1996**, 75 (13), 1533-1536.
- [19] C W Lee, P M Lemieux, B K Gullet, J V Ryan, J D Kilgroe, *Stud Environ Sci*, **1998**, 361-378.
- [20] A Khanfekr, K Arzani, A Nemati, M Hosseini, *Int. J. Environ. Sci Tech*, **2009**, 6 (1), 105-112.
- [21] P Arulazhagan, N Vasudevan, I T Yeom, *Int. J. Environ. Sci Tech*, **2010**, 7 (4), 839-852 .
- [22] E O Fagbote, E O Olanipekun, *Int J EnvironSciTech*, **2010**, 7 (3), 561-570.
- [23] L Lijklema, J M Tyson, A S Le, *Water Science and Technology*, **1993**, 12, 27.