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Chemical Profiling of Heroin Recovered from the North Korean Merchant Vessel Pong Su

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ABSTRACT: Heroin samples, seized from the North Korean merchant vessel Pong Su in Australian waters, were analyzed to determine geographic origin. Duplicate samples were analyzed by the National Measurement Institute’s Australian Forensic Drug Laboratory (AFDL) and the United States Drug Enforcement Administration’s Special Testing & Research Laboratory (STRL). Alkaloid ratios were determined by both LC-DAD and CE-DAD techniques. Acid/neutral manufacturing by-products were determined by solvent extraction followed by GC-MS. Solvents, trapped in the heroin particles during manufacture were detected by both static headspace GC-MS and purge & trap GC-MS. The alkaloid ratios obtained were consistent with heroin of a Southeast Asian origin and Principal Component Analysis of the alkaloid results demonstrated the presence of at least 4 sub-groupings within the seizure. The solvent analysis detected diethyl ether and ethyl acetate, solvents typically seen in Southeast Asian heroin. However, the acid/neutral analysis revealed compounds not normally seen in heroin of a Southeast Asian origin. Furthermore, sterol-like molecules, always detected in the acid/neutral analysis of Southeast Asian heroin, were absent from the Pong Su samples. The Pong Su heroin, although similar to Southeast Asian heroin, has sufficient differences to classify it as having an unknown origin at the time of this writing.

KEYWORDS: forensic science, chemical profiling, origin classification, heroin, alkaloids

Australia is a large continent with a coastline exceeding 35,000 kilometers and as a major trading nation it is visited each year by many merchant ships. Such an extensive coastline offers many opportunities for illicit drug importation. In April 2003 a North Korean-flagged merchant vessel passed close to the Australian coastline at Lorne, Victoria. Six packages were placed in a rubber boat for transfer to the shore. On route to shore it is alleged that one package fell overboard and its contents were never recovered. On 16th April, 2003, based on intelligence received by Australian Federal Police (AFP) officers, two packages were intercepted by federal agents from the luggage boot of a motor vehicle. On the 7th May, 2003, the remaining three packages were located by AFP agents buried near the township of Lorne. The foreign merchant vessel which had been shadowed by a Royal Australian Navy ship was eventually boarded and the crew and officers arrested (1).

Forensic deconstruction by AFP scientists revealed that of the two recovered packages from the luggage boot contained two inner packages, each containing 36 rectangular white blocks of compressed powder making a total of 144 blocks. Similarly, the 3 packages buried near Lorne contained 2 inner packages, each containing 36 rectangular blocks making a total 216 blocks. In total of 360 blocks were recovered from the 5 packages. The average weight per block was determined to be approximately 350 grams. Initial field tests indicated the material was heroin. Each block was individually wrapped with plastic wrapping material stamped with the ‘Double UO GLOBE’ logo. One hundred blocks were randomly selected and one gram (1g) core samples from each of the
100 suspected heroin blocks were sent to the National Measurement Institute’s Australian Forensic Drug Laboratory (AFDL) for identification and purity analysis. Because of the nature and size of the seizure it was decided to carry out chemical profiling on the samples to determine geographical origin. Determination of the major alkaloids, acid/neutral manufacturing by-products and occluded solvents in heroin has been used by law enforcement agencies to determine geographic origin and drug trafficking routes for many years. A knowledge of the ratios of the alkaloid impurities present in illicit heroin to the total morphine content of the sample allows the chemist to assign origin (2, 3), i.e. Southeast Asian (SEA), Southwest Asian (SWA), South American (SA) or Mexican (Mex). This assignment is achieved by comparing these alkaloid ratios with the same ratios obtained on a large number of authentic samples. A knowledge of the occluded solvents provides further evidentiary material for this rough geographic assignment. Solvents used in the processing of opium through to heroin can be indicative of the geographic origin (4). A further aid to the assignment of origin is afforded by the analysis of the acidic and neutral molecules contained within the heroin. During the transformation of opium to heroin a large number of by-products are formed. The acetylation of morphine to diacetylmorphine involves drastic reaction conditions and other co-extracted alkaloids such as thebaine, papaverine and noscapine undergo chemical reaction. Important among the by-products are various N-acetyl derivatives of noscapine, thebaine, morphine, papaverine and codeine (5). The chromatographic impurity pattern afforded by these acidic and neutral compounds has proven useful in profiling heroin (2, 5, 6). The method used by the United States DEA’s Special Testing and Research Laboratory (STRL) based on work by Allen et.al (5) has been successfully applied to heroin profiling in a number of laboratories. It is achieved by partitioning the neutral and acidic by-products from the basic heroin sample matrix containing the major alkaloids by using an acid/organic solvent mixture.

Most heroin seizures made by AFP officers at the Australian border are of Southeast Asian origin and principally involve Burma, Laos and Thailand (7). Other sources of heroin in the world are Southwest Asia, which today refers mainly to Afghanistan, and South America and Mexico. Analyses of the major alkaloids, acid/neutral manufacturing by-products and occluded solvents were performed on the Pong Su heroin samples. Data so acquired were added to Australian Illicit Drug Intelligence Program’s (AIDIP) database. Duplicate samples were sent to the Special Testing and Research Laboratory (STRL) of the United States Drug Enforcement Administration.
Experimental Section

Reagents and Standards
All reference standards, internal standards and surrogate standards used in the chemical profiling at AFDL were obtained from the reference collection of the National Measurement Institute. Bis-trimethylsilyltrifluoroacetamide (BSTFA) was obtained from Progen Biosciences, Archerfield, Qld. Analytical grade methanol was obtained from Malinckrodt Chemicals and analytical grade dichloromethane and acetonitrile from Merck, Kilsyth, Victoria. Hexylamine, propiophenone and benzpinacolone were obtained from Aldrich, Castle Hill, NSW and were used without further purification.

Major Alkaloid Analysis by Liquid Chromatography at AFDL
The heroin samples were analyzed for the alkaloids noscapine, papaverine, acetylcodine, codeine, morphine, 6-monoacetylmorphine, 3-monoacetylmorphine and diacetylmorphine by the method of Lurie et.al (8). Internal standard solutions were prepared by accurately weighing 50 mg of propiophenone into a 100 mL volumetric flask and diluting to volume with mobile phase. Each heroin sample was homogenized by lightly crushing and mixing in a mortar and pestle. 15-20 mg of sample were accurately weighed into a 10 mL volumetric flask, internal standard (1 mL) added and the whole diluted to volume with mobile phase and transferred to an injection vial. Solutions of morphine, codeine, noscapine, 6-MAM and 3-MAM (5, 10, 25, 50 and 100 mg/L), acetylcodine (10, 50, 100, 150 and 200 mg/L), heroin (50, 100, 250, 500 and 1000 mg/L), papaverine (1, 5, 10, 25 and 50 mg/L) for calibration were prepared from a stock solution made by weighing 10 mg of parent compound into 10 mL volumetric flasks and diluting to volume with mobile phase. Appropriate aliquots and internal standard (1 mL) were diluted accordingly to give the calibrator solutions. Chromatography was performed using a HP-1090 liquid chromatograph equipped with a 150mm x 3.2mm x 5µm Alltima C_{18}-LL column. The flow rate was optimized at 0.75mL/min. The mobile phases were methanol (A) and an amine-phosphate buffer (B). The amine-phosphate buffer was prepared by diluting 30mL of 2N NaOH, 11.5mL of 85% phosphoric acid and 3.5mL of hexylamine to 1 litre with Milli-Q water and filtering. The gradient profile commenced at 5% A and 95% B and changed linearly to 30% A and 70% B in 20 min, held for 6 min and changed linearly to 80% A and 20% B in 10 min, held for 4 min and then allowed to return to 5% A and 95% B. The injection volume was 20µL and the detection wavelength was 240nm.

Major Alkaloid Analysis by Capillary Electrophoresis at STRL
Samples were analyzed for heroin and the major alkaloids using capillary zone electrophoresis with dynamically coated capillaries by the method of Lurie et. al. (9). Standards, except for O6-monoacetylmorphine hydrochloride, which was prepared separately, were prepared as a mixture in the injection solvent which was a 2:8 mixture of methanol and 3.75mM phosphate buffer (pH 3.2) (9). Each sample was weighed out as a 20 mg-equivalent of heroin HCl (based on separate analytical results), placed into a 50 mL
volumetric flask (final concentration 0.4 mg/mL) and diluted to volume with injection solvent. After a 15 minute sonication, the preparation was filtered and 1mL added to a 2 mL glass CE vial and analyzed on an Agilent CE Model #G1600AX using a HPCE standard capillary 50 µm id. and following the method as described by Lurie et. al. (9).

Solvent Analysis by Purge & Trap GC/MS at AFDL
Occluded solvents were qualitatively determined by purge and trap GC/MS by a method used by the National Measurement Institute’s Trace Organics laboratory (10). Each heroin sample (20 mg) was weighed into a headspace vial which was then filled with a 2% aqueous solution of sodium sulfate and capped. The vial was submitted for solvent analysis using a Tekmar 3000 Purge & Trap integrated with an Agilent 6890/5973 GC-MS (5). The purge rate was 40mL/min for 11 minutes followed by a dry purge for 3 minutes and desorption at 260°C for 2 minutes. The transfer line to the GC was held at 100°C. The GC system was fitted with a 30 m x 0.25 mm i.d. x 0.25 µm film thickness DB-624 column. The oven temperature was programmed as follows: initial temperature100°C (4 min) and ramped to 120°C at 7°C/min (0 min) and ramped to 220°C at 15°C/min (2 min). The injector was operated in the split mode at 180°C with a split vent flow of 15mL/min at 140kPa. The MSD was operated in the electron ionization mode (EI) at 70eV and a scan range of 34 to 280. The solvent delay was 1.2 min and the transfer line to the MSD was held at 245°C. The method identified the presence or absence of solvents, including diethyl ether, ethyl acetate, acetone, chloroform, dichloromethane, toluene, benzene and the xylenes, acetonitrile.

Solvent Analysis by Headspace GC/MS at STRL
Occluded solvent analysis for each sample was determined using a static headspace-gas chromatography-mass spectrometry method (4). A 40 mg heroin equivalent for each sample was weighed into a headspace vial. Aqueous sodium sulfate (22%) containing five deuterated internal standards was added to each sample and to each of the low, mid and high calibration standard solutions. Solvent analysis was performed using static headspace-gas chromatography-mass spectrometry (Agilent 7964, Agilent 6890/5973 GC-MS) according to the method of Morello and Meyers (4).

Acid/Neutral Analysis by GC/MS at AFDL
Acid/Neutral manufacturing by-products were analyzed using a modification of the method used by STRL as described by Allen et. al. (5) and Neumann and Gloger (2). Samples were prepared by weighing 30 mg equivalents (based on the total morphine results obtained from the alkaloid analysis) into 1 mL conical tubes and dissolving in 4 mL of light petroleum (b.p. 40-60 °C)/dichloromethane mixture (60/40). Internal standards, 2,2,4-trimethylacetophenone (50 mg/L; 100 µL) and d₉-O6,O3,N-triacetylnormorphine (50 mg/L; 100 µL) and the surrogate standard N-propionylmorphine (50 mg/L; 100 µL) were added to each sample, reagent blank and QA sample. Sulfuric acid (4 mL, 0.25 M) was then added to each tube and the sample placed on a rock-and-roll mixer for 10 minutes. The upper organic layer (3 mL) was removed and concentrated just to dryness under a gentle flow of dry nitrogen in a reacti-vial. The residue was immediately dissolved in 250 µL of a mixture of BSTFA/hexane (50/50), capped and heated at 70 °C for 30 minutes. After cooling 100
μL were transferred to a limited insert GC vial. An Agilent 6890/5973 GC/MSD equipped with a 30 m x 0.25 mm i.d. DB-5MS column with 0.25 μm film thickness was used. The column temperature was programmed from 100 °C (1 min) to 240 °C at 6 °C/min and then to 280 °C at 2 °C/min and finally to 320 °C at 6 °C/min. The injection port temperature was 280 °C operated in the splitless mode for 0.3 minutes. The MSD was operated in the selected ion monitoring mode. Compounds monitored and their target ions are given in Table (1).

After data acquisition, peak areas of the analytes of interest (Table 1) were uploaded automatically as CRD files using the Agilent ChemStation software into the database. The peak area of each analyte was summed and each peak area then expressed as a percentage of the total area. The ratio of the peak area of each analyte to the peak area of the internal standard, and the recovery of the surrogate standard were also determined.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Target Ions</th>
</tr>
</thead>
<tbody>
<tr>
<td>3,4-dimethoxy-4,5-epoxyphenanthrene</td>
<td>194(100), 253(65)</td>
</tr>
<tr>
<td>thebaol (O4-TMS)</td>
<td>296(100), 326(74)</td>
</tr>
<tr>
<td>acetylthebaol</td>
<td>254(100), 239(73), 296(35)</td>
</tr>
<tr>
<td>4-acetoxy-3,6-dimethoxy-5-[2(NMA)]ethylphenanthrene</td>
<td>265(100), 252(75), 395(40)</td>
</tr>
<tr>
<td>O6,O3,N-triacetylnormorphine</td>
<td>209(100), 87(95)</td>
</tr>
<tr>
<td>d0- O6,O3,N-triacetylnormorphine (ISTD)</td>
<td>210(100), 87(96)</td>
</tr>
<tr>
<td>O6,N-diacetylnormorphine</td>
<td>266(100), 281(61), 87(54)</td>
</tr>
<tr>
<td>N-acetylnorlaudanosine</td>
<td>234(100), 192(82)</td>
</tr>
<tr>
<td>N-propionynorlaudanosine (ISTD)</td>
<td>192(100), 248(95)</td>
</tr>
<tr>
<td>papaverine</td>
<td>338(100), 324(95), 308(36)</td>
</tr>
<tr>
<td>noscapine</td>
<td>220(100), 205(12)</td>
</tr>
<tr>
<td>N-acetylnonoscapine</td>
<td>248(100), 206(77), 191(28)</td>
</tr>
<tr>
<td>O6,N-diacetylnorcodeine</td>
<td>223(100), 369(52), 87(64)</td>
</tr>
<tr>
<td>4-acetoxy-3,6-dimethoxy-8-[2(NMA)]ethylphenanthrene</td>
<td>280(100), 267(30), 395(35)</td>
</tr>
<tr>
<td>(E)-N-acetylanhydrornornarcine</td>
<td>382(100), 193(98), 455(16)</td>
</tr>
<tr>
<td>(Z)-N-acetylanhydrornornarcine</td>
<td>382(100), 193(98), 455(21)</td>
</tr>
<tr>
<td>(1R,9S)-1-acetoxy-N-acetyl-dihydroanhydrornornarcine</td>
<td>280(100), 252(42)</td>
</tr>
</tbody>
</table>

**Acid/Neutral Analysis by GC/MS at STRL**

Acidic and neutral manufacturing impurities in heroin samples were isolated and analyzed by Gas Chromatography-Mass Spectrometry employing a method originally done using gas chromatography-flame ionization (2, 6). A 45 mg morphine equivalent of each sample was placed into a centrifuge tube and dissolved in 5mL of a mixture prepared from petroleum ether (b.p. 20-40°C) (540mL) and methylene chloride (360mL). 2.0N Sulfuric acid (4mL), containing 10% sodium sulfate, was added to each sample and vortexed to extract the acid-neutral fraction into the organic phase. Following
centrifugation the organic phase was isolated, concentrated to dryness and the residue was derivatized with MSTFA. The derivatized extract was analysed for acidic and neutral manufacturing by-products on a Polaris-Q GC-MS system in the full scan mode ($m/z$ 100-575). The initial injector temperature was 85$^\circ$C (1.5 minute hold) and ramped to 295$^\circ$C (24 min hold) at 180$^\circ$C/minute. Injection mode was splitless. The initial oven temperature was 60$^\circ$C (6 minutes hold time), ramped to 200$^\circ$C at 40$^\circ$C/minute followed by a second ramp to 250$^\circ$C at 8$^\circ$C/minute and a third ramp to 295$^\circ$C (11 minutes hold time) at 1.5$^\circ$C/minute. The carrier gas was helium at a constant velocity of 52.0 cm/second

**Statistical and Chemometric Analysis of Data**
The existence of sub-groups within the seizure, and its relation to other Australian seizures of Southeast Asian heroin, were of interest. The ratios total codeine/total morphine, noscapine/total morphine and papaverine/total morphine are used by the UN and DEA (11) to assign the general origin of a sample, and graphs of these provide a useful way of identifying groups within a seizure. Some benefit may be gained from a study of the principal components of the data. A principal component is a linear combination of the data that contains the maximum variance, and so should help to distinguish sub-groups within the data. A principal components analysis (PCA) on the raw data of acetylcodeine, heroin, 6-MAM and noscapine for the 100 samples was performed with mean centering but not standardization, using Matlab (v 7.0, Mathworks Inc, USA) and Excel (Office 2002, Microsoft, Seattle) with the add-in XLStat (v7.5, Addinsoft, USA).

**Results and Discussion**

**Alkaloid Profiling**
The composition of the major alkaloids in the 100 seized samples analyzed in the AFDL are summarized in Table 2. The bracketed values are the results obtained by STRL.

<table>
<thead>
<tr>
<th></th>
<th>6-MAM %</th>
<th>Diacetylmorphine %</th>
<th>Acetyl codeine %</th>
<th>Noscapine %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Median</td>
<td>0.3 (0.3)</td>
<td>66.0 ( 67.2)</td>
<td>16.4 (16.5)</td>
<td>0.3 (0.2)</td>
</tr>
</tbody>
</table>
Table 3 summarizes the ratio of major alkaloids to total morphine obtained from these 100 samples. Again the bracketed figures are the ratios obtained by STRL. The results obtained by the two laboratories agree well. Using either set of ratios, and comparing these ratios with those obtained for heroin samples of known origin (‘authentics’) all the samples analyzed would be categorized as being of a Southeast Asian origin (3). This is noteworthy because although the same heroin blocks were examined by both national laboratories, different core samples from these blocks had to be used for the origin determination in each laboratory.

TABLE 3- Major alkaloid to total morphine ratios.

<table>
<thead>
<tr>
<th></th>
<th>Codeine/Morphine</th>
<th>6-MAM/Morphine</th>
<th>Noscapine/Morphine</th>
</tr>
</thead>
<tbody>
<tr>
<td>Median</td>
<td>27.1 (24.1)</td>
<td>0.5 (0.5)</td>
<td>0.5 (0.3)</td>
</tr>
<tr>
<td>Minimum</td>
<td>12.5 (10.7)</td>
<td>0.3 (0.2)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Maximum</td>
<td>52.1 (49.0)</td>
<td>1.0 (0.9)</td>
<td>2.1 (1.5)</td>
</tr>
</tbody>
</table>

However, an inspection of the data reveals some interesting results. A large number of samples contained unusually high acetylcodeine content with 17 samples having acetylcodeine levels between 24.7% and 27.2%. Many samples contained higher levels of noscapine than is typical for Southeast Asian heroin. While 18 samples contained no noscapine at all, 15 contained noscapine at levels between 0.6% and 0.9%. Figure 1 shows the noscapine/morphine ratio plotted against the codeine/morphine ratio for all samples. The groupings that appear in the plot of Figure 1 may be enhanced by inspection of the principal components. PC1, which explains 98% of the variance, and has a positive loading for heroin and negative loading for acetylcodeine, when plotted in ascending order, shows at least four clear groups (Figure 2).
All heroin samples received by the AFDL during the years 2000–2003 were profiled for geographic origin. It was determined that of these, 1533 samples were of a Southeast Asian origin. A comparison of acetylcodine and noscapine levels measured in the heroin samples seized from the Pong Su with the 1533 samples classified as Southeast Asian is given in Figure 3. It is apparent that although the Pong Su samples may be classified as SEA based on their alkaloid ratios some of them are not typical of heroin traditionally produced in this region. While a number of the Pong Su samples fall in the body of the historical heroin SEA seizures, there is a clear group with high codeine and non-zero noscapine that is not typical of earlier analyses.
The solvent analysis performed at both AFDL and the DEA revealed the presence of diethyl ether and ethyl acetate in each heroin sample seized from the Pong Su. This solvent profile has been recognised by chemists working at the DEA’s Special Testing and Research Laboratory as being typical of Southeast Asian heroin (4). Occluded solvents found in heroin samples are indicative of processes used to prepare heroin. Heroin prepared in Southwest Asia typically contains acetone, while heroin produced in Mexico and South America may contain a variety of solvents including methyl ethyl ketone, methyl isobutyl ketone, acetonitrile, acetic acid and xylenes. Heroin samples examined at AFDL and determined to be from Southeast Asia on the basis of their alkaloid ratios all contained diethyl ether and ethyl acetate.
FIG. 3- Data of figure 1 (crosses) with 1533 samples of authentic Southeast Asian heroin seized in Australia during 2002-2003 (closed circles). The group within the dashed oval is that identified as “1” in the PC plot of Figure 2.

FIG. 4- Total ion current chromatogram of the acid/neutral extract of a heroin sample seized from the Pong Su. 1: O6,N-diacetylnorcodeine; 2: O6,N-
The analysis of the acidic and neutral components by both laboratories revealed the presence of the following manufacturing by-products in each sample: O6,N-diacetylnorcodeine, O6,N-diacetylnormorphine and O3,O6,N-triacetylnormorphine. A chromatogram showing the presence of these three compounds in one of the samples seized from the Pong Su is presented in Figure 4. These three compounds are frequently detected in heroin samples determined, by their alkaloid ratios, to be of Southeast Asian origin. They are usually the most abundant N-acetylated by-products found in heroin produced in this region. A chromatogram showing the acidic and neutral manufacturing by-products in an authentic Southeast Asian heroin sample is given in Figure 5.

![Chromatogram showing the presence of three compounds](image)

**FIG. 5-** Total ion current chromatogram of the acid/neutral extract of a typical SEA heroin sample seized in Australia. 1: O6,N-diacetylnorcodeine; 2: O6,N-diacetylnormorphine; 3: O6,O3,N-triacetylnormorphine.

The acid/neutral analysis of each of the seized samples also revealed the presence of 4-acetoxy-3,6-dimethoxy-5-[2-N-methylacetamido]ethylphenanthrene. This compound is an N-acetylated degradation product of the opium alkaloid thebaine. Also detected in each sample was noscapine which is an N-acetylated degradation compound of noscapine. A chromatogram showing these compounds is given in Figure 4. These manufacturing by-products are not typically seen in heroin of Southeast Asian origin but are observed in heroin of Southwest Asian origin.

Also of interest was the absence of peaks in the acid/neutral chromatograms that are believed to be due to a group of unidentified sterol-like molecules. The STRL has
tentatively identified the presence of these molecules, only observed to date in heroin of known Southeast Asian origin, as sterols. While the exact structures of these sterols are as yet unknown, their mass spectral fragmentation patterns are consistent with molecules having the steroid nucleus. The chromatographic peaks of this group of compounds, found in an earlier seizure from Southeast Asia can be seen in Figure 5. A mass spectrum of one of these compounds, tentatively identified as a sterol molecule is shown in Figure 6. These sterols were not detected by either AFDL or STRL in the samples seized from the Pong Su. Both national laboratories have detected these sterol-like molecules in all authentic Southeast Asian heroin samples examined to date. The absence of these compounds in the heroin seized from the Pong Su is noteworthy.

FIG. 6- Mass spectrum tentatively identified as being due to a member of a group of sterol-like molecules detected only in heroin of a Southeast Asian origin.

Two possible explanations may be that (1) opium of Southwest Asian origin is being transported into Southeast Asia for processing into heroin or; (2) a new form of heroin chemically similar to the Southeast Asian heroin and processed in a very similar way has been produced. Evidence for the latter suggestion is the observation of an increasing noscapine concentration in SEA heroin in Australia over the last 4 years. One possible explanation for the variation within a single shipment is the theory of central collection, pooling and distribution points for either the heroin or the opium.
Conclusions
Despite having major alkaloid and occluded solvent profiles consistent with a Southeast Asian origin, the heroin seized from the North Korean merchant vessel ‘Pong Su’ was classified as of unknown origin. This decision is based on the presence of compounds in the acid/neutral analysis normally not seen in heroin from this region and the absence of sterols that are normally present in Southeast Asian heroin. The presence of acid/neutral manufacturing by-products common to heroin of Southwest Asian origin, in heroin samples with an otherwise Southeast Asian profile is at present inexplicable. Future heroin seizures will be examined for evidence of a similar profile.

Further work, focussing on the carbon stable isotope ratios of authentic heroin samples from Southeast Asia, Southwest Asia, South America and Mexico and the Pong Su samples, has been carried out at the STRL (12). This work and the alkaloid analyses has demonstrated that the Pong Su samples are different from any heroin previously encountered. However, the geographic origin of this seizure will remain unknown until an authentic heroin sample with a matching profile is obtained.

Acknowledgements
The authors are indebted to the Australian Federal Police for providing the samples.
References