CALL FOR NOMINATIONS

It’s time for elections for IAOS President. Deadline for nominations is November 1, 2008. Nominations may be sent via email to current IAOS President, Anastasia Steffen at asteffen@unm.edu.

CONSIDER PUBLISHING IN THE IAOS BULLETIN

The Bulletin is a twice-yearly publication that reaches a wide audience in the obsidian community. Please review your research notes and consider submitting an article, research update, or lab report for publication in the IAOS Bulletin. Articles and inquiries can be sent to cdillian@princeton.edu. Thank you for your help and support!

DELPHI, GREECE, OBSIDIAN CONFERENCE: January 11-14, 2008
The Dating and Provenance of Natural and Manufactured Glasses

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CONFERENCE ABSTRACTS ARE NOW AVAILABLE ON THE IAOS WEBSITE:
www.peak.org/obsidian
NOTES FROM THE PRESIDENT

The Society for American Archaeology annual meetings in Vancouver, B.C., were busy with IAOS activities this past March. These included a symposium organized by Rob Tykot with papers reporting obsidian sourcing techniques and research in a wide diversity of locations across several continents. The IAOS Workshop on Sourcing and Dating of Obsidian, organized by Chris Stevenson and Michael Glascock, provided an opportunity for review of current methods and techniques for both obsidian sourcing and hydration analyses. The demonstration of a portable XRF spectrometer was particularly interesting and provided an opportunity for discussion of the pros and cons in using these increasingly popular instruments. Submissions for next year’s SAA Meetings in Atlanta will be due in early September. If you are interested in organizing symposia or other events suitable for IAOS sponsorship at these or other professional meetings or workshops, I encourage you to contact me.

Good attendance at the IAOS Annual Meeting allowed for productive discussion of use of the IAOS membership contact list, the status of IAOS officer positions, and proposed changes to the IAOS By-laws. It was agreed that current Secretary-Treasurer Colby Phillips will continue for another two-year term (which requires a change to the By-laws, as discussed below). Two nominations were received for President; additional nominations can be sent to me or the Secretary-Treasurer (to be received by October 1, 2008). Several nominations for the IAOS student poster/paper award were submitted. Most of the nominations were for presentations at the 2008 SAA Meetings. If you know of an excellent student research paper/poster from other conferences this year, contact me so that IAOS can acknowledge their work.

Proposed changes to the IOAS By-laws were prompted during review by the Executive Board this past winter. Several changes are proposed based on this review and subsequent discussion at the Business Meeting. To summarize, we are proposing the elimination of Institutional Member status, change in membership year to calendar year, change in language throughout the By-laws to reflect the use of email as the primary means of communication with the membership, specification of how member contact information will be used, and change in the Secretary-Treasurer office to allow consecutive terms for this position. A copy of the By-laws showing the current text and proposed changes will be posted on the IAOS website at www.peak.org/obsidian. Please review the proposed changes and send your comments (to myself or the Secretary-Treasurer) by September 1, 2008. Once member comments are received, a final amended version will be available for a vote by the membership either through email or at the next Business Meeting. We are especially interested in comments from longtime IAOS members on the proposed elimination of the Institutional Member category. While this was a substantive part of the early structure of IAOS, there currently are no active Institutional Members. In reviewing the By-laws, let us know if this category should be eliminated or retained and revitalized.

Ana Steffen
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IAOS Membership Update

The International Association for Obsidian Studies is in the process of transitioning all IAOS memberships to a Jan. 1 - Dec. 31 calendar year. If you joined or renewed your membership any time last year before Oct. 1 2007, please renew your membership now (you can renew online via PayPal on the IAOS website at http://www.peak.org/~obsidian or simply use the form on the last page of this Bulletin). Renewing your membership will allow the IAOS to continue. If you are unsure about your membership anniversary or last renewal date, please send a message to me at colbyp@u.washington.edu and I will check it for you.

Regards,
Colby Phillips, IAOS Secretary/Treasurer
Field Exploration and Instrumental Neutron Activation Analysis of the Obsidian Sources in Southern Armenia

John F. Cherry, Brown University
Elissa Z. Faro, Brown University
Leah Minc, Oregon State University

Introduction

The Armenian highlands are an area of intense tectonic activity: within Armenia alone, some 450 volcanic domes have been recorded, and there are as many as 26 reported obsidian outcroppings in the southern Caucasus spread across five Plio-Pleistocene volcanic regions (Blackman et al. 1998: Table 1). Compared to the Mediterranean, Anatolia, and certain other parts of the Near East, field exploration and geochemical characterization of the Armenian obsidian sources has been slow to develop. Since the 1990s, however, much new work has been undertaken — most notably, a comprehensive program of sourcing and provenience studies by Dr. Ruben Badalyan and colleagues, involving the instrumental neutron activation analysis (NAA) of geological hand samples from the majority of the sources and of a further 576 samples derived from 53 archaeological sites throughout the region (Badalyan 2002; cf. Oddone et al. 2000). The most southerly Armenian sources in the Syunik region (at Satanakar, Metz Sevkar, Pokr Sevkar, and Bazenk), however, have not yet been adequately characterized by any of the obsidian research programs of the last two decades (see, e.g., Keller et al. 1994: Table 4, listing analyses of just nine samples). This is unfortunate, since a much earlier study using NAA (Renfrew and Dixon 1977: 145 and Tables 1-2) had suggested that obsidian artifacts of their “Group 3c”, all from sites in Iranian Azerbaijan, probably derived from an unlocated Armenian source in, or to the north of, the Lake Urmia region; the Syunik sources (unknown at the time to these authors) are the only ones that come close to matching this description.

The inception in 2005 of the Vorotan Project — a collaborative Armenian-American program of archaeological survey and excavation in the middle reaches of the Vorotan River in Syunik province — has provided an opportunity to conduct more thorough exploration of the Syunik sources, which lie ca. 30 km northwest of the project’s main area of focus (Figure 1). Within this latter area, stratified excavation, systematic gridded surface collection of sites, and intensive pedestrian survey conducted by the project have so far yielded a total assemblage of 8,690 pieces of obsidian, whose most likely origin is one or more of the Satanakar, Sevkar, and Bazenk flows. Study has concentrated on documentation and sampling of these flows, as well as secondary obsidian deposits in the Vorotan River, together with morphological, technological, and metrical studies of the assemblages collected by survey and excavation.

Figure 1. Map showing the location of the southern Armenian Syunik obsidian sources, in relation to the area under study by the Vorotan Project and sites from which obsidian artifacts have been analyzed by INAA. (Map created by Lynn Carlson and Elissa Faro.)
The Syunik Obsidian Flows

The Syunik obsidian flows lie close to the Armenian border with Azerbaijan, in remote and rugged mountainous uplands reaching elevations above 3,200 m. Fieldwork was conducted at all of the geological sources in 2006, with supplementary visits to collect additional samples in 2007. Since these liparite domed volcanoes have already been well described from the geological standpoint (Karapetyan 1972: 73-80), the goals were to locate the various obsidian flows accurately using handheld Trimble GPS/GIS units, to document them photographically, to make systematic collections of geological hand-samples for NAA, and to inspect the evidence for knapping workshops noted, but not described, by earlier visitors (Badalyan et al. 2004: 455-456).

The obsidian occurs within rhyolitic and perlitic flows on the flanks of these volcanoes. It is in general dark gray to jet black, although at the Sevkar sources, uniquely, there are significant occurrences of red-brown mottling; it is translucent when thinly flaked, and for the most part free of perlitzation or other crystalline structures that would limit its suitability for knapping. Raw material is available in large blocks between 25 and 75 cm in diameter (sometimes even larger). The abundance of these flows is so great that they can be readily discerned in imagery from space; in one location, on the south flanks of Satanakar, the obsidian outcrops in the form of a spectacular cliff exceeding 50 m in height, with a vast talus slope below it providing a ready source of raw material in convenient form (Figure 2). Indications of knapping activities at the source itself are frequent at Metz and Pokr Sevkar, especially on the slopes of the latter, where macrocores, extremely large core-trimming flakes, portions of blade-cores, blade-flakes, and even large segments of finished blades have been recorded.

Sampling and Analytical Procedures

At least a dozen hand-samples were collected from each source, at regular intervals across as much of the flow as possible, and including examples that reflected macroscopically visible differences in the material (e.g., in terms of color, presence of phenocrysts, etc.). A total of 66 fist-sized geological samples collected in 2006-7 were submitted for trace-element analysis via Instrumental Neutron Activation Analysis (INAA) at the Oregon State University Radiation Center. In addition, 69 pieces of artifactual material from sites within the Vorotan Project study area were included in this analytical program: they come from Chalcolithic, Middle Bronze Age, and Iron Age III (“Yervandid”) contexts at the sites of Nerkin Godedzor, Shaghat 1, Shaghat 2, and Shaghat 3.
and Aghidu (see Figure 1). Twelve river-rolled cobbles, collected both from the banks of the River Vorotan and during surface survey activities, make up the remainder of the samples.

The obsidian samples were analyzed for a suite of 35 major, minor, and trace elements, through a sequence of two separate irradiations and multiple counts of resultant gamma activity. These two counts provided data on As, Br, La, Lu, Mo, K, Na, Sm, U, Yb, and Ba, Ce, Co, Cr, Cs, Eu, Fe, Hf, Nd, Rb, Sb, Sc, Sr, Ta, Tb, Th, Zn, and Zr, respectively. Elements with short and intermediate half-lives, by contrast, were analyzed using the pneumatic tube irradiation system, which provided data on Al, Ca, Cl, Dy, Mn, K, Na, Ti, and V. Full details of the procedures employed will be presented elsewhere.

Results

The INAA results obtained for the geological samples in this study have been evaluated through a series of bivariate plots of element concentrations, cluster analysis, and principal components analysis. It is apparent that the Bazenk, Satanakar, and Sevkar sources can be clearly discriminated on the basis of their concentrations of the lighter rare earth elements (especially La and Ce) and Th. The separation of these sources is illustrated in Figure 3, which uses the elements La and Th; ellipses around the subgroups are plotted at the 95% confidence level. Samples from Pokr Sevkar and from widely separate areas of the Metz Sevkar source display generally overlapping elemental compositions, and it is possible to treat these flows together as the “Greater Sevkar” complex, which itself constitutes a fairly tight group. Satanakar has provided trace elemental evidence of two quite clearly distinguishable flows, which also map onto the volcano in a spatially coherent manner.

The assignment of obsidian artifacts to their most likely geological source shows a strong preference for the Greater Sevkar flows (Figure 4). This is an unsurprising result, given that they are geographically very extensive, at lower altitude, and more readily accessible than the other two sources. Artifactual samples from late Chalcolithic Godedzor (made available to us courtesy of Dr. Pavel Avetisyan) are dominantly of Sevkar obsidian, but with minor use of the Satanakar and Bazenk sources too, and no evidence for exploitation of any other Armenian obsidian source. All the samples from Shaghat 1 (mainly Middle Bronze Age, but a few from mid- to late-1st millennium B.C. contexts) are attributable to the Sevkar flows, with the exception of a single artifact that appears closest in composition to published data on samples from volcanic groups in the northern part of Region III in central Armenia (Keller et al. 1994: 71, figs. 1, 3; see also Blackman et al. 1998; Oddone et al. 2000). Lastly, obsidians from the Iron Age III (mid-first millennium B.C.) citadel at Aghidu are once again dominantly from Sevkar, with a single example from Bazenk and another probably from the Kecheldag/Kel’badzhar source, which lies just across the border in Azerbaijan (and thus could not be included in the present program of fieldwork).
Conclusions

It comes as little surprise that the occupants of sites only a few dozen kilometers from the high-quality Syunik obsidian sources should have relied on them almost exclusively. On the other hand, since they occur at such high altitude, these sources would have been inaccessible for more than half the year, and it is likely that the rolled obsidian cobbles available as a secondary deposit along the River Vorotan constituted a raw material resource equal in importance to the flows in situ. Now that the Syunik sources have been much more thoroughly characterized than hitherto, the logical next step is a comprehensive program of analysis of obsidian artifacts from Azerbaijan, northwestern Iran, Nakichevan, and southeastern Anatolia, in order to establish the regional scale of the exploitation of these sources (and whether they have any connection to the “3c source” identified long ago by Renfrew and colleagues). We also intend to explore the metrical aspects of the archaeological assemblage in greater detail, as well as seeking an explanation for a dramatic decline in the abundance of obsidian artifacts after the Middle Bronze Age.

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Renfrew, C., and J. Dixon
Introduction

The discovery of obsidian on archaeological sites in eastern Anatolia and the surrounding regions shows that obsidian utilization took place over a long period of time from at least the early Neolithic through Bronze Age. Renfrew et al. (1966; 1968) demonstrated that the chemical compositions of obsidian artifacts could be used to reconstruct obsidian exchange networks involving the Anatolian sources. As a result, archaeologists have subdivided the ancient Near East into regions according to the particular source(s) utilized. A few of the Near Eastern obsidian sources were associated with long-distance exchange networks, but many obsidians did not travel far from their origin.

Background

Studies of obsidian in Iran have not progressed as rapidly as in other regions resulting in an imperfect model for regional source utilization. Using XRF and NAA to characterize obsidian artifacts from the archaeological sites at Salmas and Kaleybar in NW Iran (see Figure 1), new information suggests the possible existence of obsidian sources in NW Iran (candidate locations for obsidian sources are Mt. Sabalan and Mt. Sahand). Models of obsidian exchange in Iran should therefore consider these “new” sources along with currently known obsidian sources in the Lake Van region of eastern Turkey and in the countries of Armenia and Azerbaijan (Blackman et al. 1998; Chataigner et al. 1998; Keller et al. 1996). The obsidian sources already known to this region are shown in Figure 1.

Archaeological Sites of Salmas and Kaleybar

The Salmas Plain (1560 m above sea level) is located in the West Azerbaijan province of Iran about 20 km west of Lake Urmia and 35 km east of the border with Turkey. The Salmas Plain was occupied continuously from the Neolithic to the Bronze Age. Evidence indicates that the earliest
inhabitants of Salmas were primarily agricultural and they used tools made of bone, ground stone, and obsidian as early as the the 6th millennium. Approximately 200 obsidian artifacts with colors primarily black and grey, have been discovered at sites in the Salmas Plain (Kargar 1996) indicating that the occupants had access to obsidian sources. The nearest sources to Salmas are located in the Lake Van region with distances ranging from 150-200 km. A total of seven obsidian artifacts from the Salmas region were made available for compositional analysis by Mr. Kargar.

The Kaleybar sites are located in the Kaleybar township of the East Azerbaijan province a distance of about 110 km northeast of the city of Tabriz and 40 km from the Aras River which serves as the border between Iran and the country of Azerbaijan. Numerous settlements from the Chalcolithic and Urartu periods are found throughout Kaleybar. Obsidian artifacts are found in greater abundance in the Kaleybar regions indicating that access to obsidian sources was probably greater (Feizkhah 2007). The nearest known sources are located in Armenia and Azerbaijan, but it is possible that cobbles of obsidian were extracted from the Aras River or from yet to be discovered sources in Iran. Mr. Feizkhah made a total of 38 obsidian artifacts from Kaleybar available for geochemical analysis.

**Analytical Methodology**

A combination of the two most reliable analytical techniques for obsidian characterization – X-ray fluorescence (XRF) and neutron activation analysis (NAA) – were employed in this investigation using the facilities available in the Archaeometry Laboratory of the University of Missouri Research Reactor Center. The advantages of XRF and NAA are well known. In short, XRF is non-destructive, more rapid, and less expensive; however, XRF measures fewer elements, has less precision, and is less accurate than NAA. By employing both methods to our study, we were able to maximize the advantages of both techniques. The entire collection of 45 of the obsidian artifacts were analyzed by XRF in order to sort the artifacts into specific geochemical types. Afterwards, a sub-sample consisting of 22 artifacts, selected to include all possible varieties, were characterized by NAA to obtain high precision data and allow direct comparison to obsidian source data reported earlier by Blackman et al. (1998) and Keller et al. (1996).

**Figure 2a.** Plot of Iron versus Europium showing NAA data for obsidian artifacts from Salmas projected against 95% confidence ellipses for obsidian sources in the Lake Van region. The Nemrut source has four subgroups. The unknown sources do not match any of the published data for obsidian sources in Turkey or the Transcaucasus region.

**Figure 2b.** Plot of Iron versus Europium showing NAA data for obsidian artifacts from Kaleybar projected against 95% confidence ellipses for obsidian sources in the Lake Van region. The Nemrut source has four subgroups.
Table 1. Summary of sourcing results for artifacts from the sites of Salmas and Kaleybar.

<table>
<thead>
<tr>
<th>Source name</th>
<th>Salmas</th>
<th>Kaleybar</th>
</tr>
</thead>
<tbody>
<tr>
<td>Meydan</td>
<td>6</td>
<td>1</td>
</tr>
<tr>
<td>Nemrut</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Unknown Group #1</td>
<td>0</td>
<td>31</td>
</tr>
<tr>
<td>Unknown Group #2</td>
<td>0</td>
<td>5</td>
</tr>
<tr>
<td>Unknown Group #3</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>7</td>
<td>38</td>
</tr>
</tbody>
</table>

Table 2. Concentrations of elements measured by NAA in groups of obsidian unknowns from Kaleybar.

<table>
<thead>
<tr>
<th>Element</th>
<th>Unknown Group #1 (n = 8)</th>
<th>Unknown Group #2 (n=5)</th>
<th>Unknown Group #3 (n=1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ba (ppm)</td>
<td>64 ± 13</td>
<td>46 ± 5</td>
<td>62</td>
</tr>
<tr>
<td>La (ppm)</td>
<td>33.5 ± 0.5</td>
<td>17.0 ± 0.5</td>
<td>27.3</td>
</tr>
<tr>
<td>Lu (ppm)</td>
<td>0.396 ± 0.012</td>
<td>0.622 ± 0.013</td>
<td>0.456</td>
</tr>
<tr>
<td>Nd (ppm)</td>
<td>12.1 ± 1.1</td>
<td>12.2 ± 0.9</td>
<td>9.8</td>
</tr>
<tr>
<td>Sm (ppm)</td>
<td>2.39 ± 0.06</td>
<td>3.93 ± 0.12</td>
<td>2.16</td>
</tr>
<tr>
<td>U (ppm)</td>
<td>10.0 ± 0.3</td>
<td>15.1 ± 0.3</td>
<td>11.7</td>
</tr>
<tr>
<td>Yb (ppm)</td>
<td>1.45 ± 0.02</td>
<td>2.39 ± 0.05</td>
<td>1.47</td>
</tr>
<tr>
<td>Ce (ppm)</td>
<td>56.2 ± 0.8</td>
<td>36.0 ± 1.4</td>
<td>45.9</td>
</tr>
<tr>
<td>Co (ppm)</td>
<td>0.130 ± 0.016</td>
<td>0.105 ± 0.047</td>
<td>0.093</td>
</tr>
<tr>
<td>Cs (ppm)</td>
<td>4.48 ± 0.08</td>
<td>7.48 ± 0.19</td>
<td>5.16</td>
</tr>
<tr>
<td>Eu (ppm)</td>
<td>0.162 ± 0.003</td>
<td>0.204 ± 0.003</td>
<td>0.111</td>
</tr>
<tr>
<td>Fe (ppm)</td>
<td>4807 ± 66</td>
<td>3612 ± 57</td>
<td>4425</td>
</tr>
<tr>
<td>Hf (ppm)</td>
<td>3.68 ± 0.05</td>
<td>3.11 ± 0.06</td>
<td>3.73</td>
</tr>
<tr>
<td>Rb (ppm)</td>
<td>173 ± 3</td>
<td>200 ± 3</td>
<td>190</td>
</tr>
<tr>
<td>Sb (ppm)</td>
<td>0.212 ± 0.006</td>
<td>0.631 ± 0.024</td>
<td>0.237</td>
</tr>
<tr>
<td>Sc (ppm)</td>
<td>1.74 ± 0.04</td>
<td>3.08 ± 0.06</td>
<td>1.97</td>
</tr>
<tr>
<td>Sr (ppm)</td>
<td>&lt; 15</td>
<td>&lt; 15</td>
<td>&lt; 15</td>
</tr>
<tr>
<td>Ta (ppm)</td>
<td>2.17 ± 0.04</td>
<td>4.49 ± 0.11</td>
<td>2.29</td>
</tr>
<tr>
<td>Tb (ppm)</td>
<td>0.188 ± 0.005</td>
<td>0.509 ± 0.019</td>
<td>0.152</td>
</tr>
<tr>
<td>Th (ppm)</td>
<td>32 ± 1</td>
<td>27 ± 1</td>
<td>35</td>
</tr>
<tr>
<td>Zn (ppm)</td>
<td>37 ± 3</td>
<td>34 ± 1</td>
<td>39</td>
</tr>
<tr>
<td>Zr (ppm)</td>
<td>165 ± 5</td>
<td>180 ± 8</td>
<td>163</td>
</tr>
<tr>
<td>Al (%)</td>
<td>6.85 ± 0.24</td>
<td>7.07 ± 0.06</td>
<td>6.76</td>
</tr>
<tr>
<td>Cl (ppm)</td>
<td>319 ± 36</td>
<td>215 ± 17</td>
<td>335</td>
</tr>
<tr>
<td>Dy (ppm)</td>
<td>1.35 ± 0.34</td>
<td>3.48 ± 0.30</td>
<td>1.49</td>
</tr>
<tr>
<td>K (%)</td>
<td>3.89 ± 0.07</td>
<td>3.70 ± 0.22</td>
<td>3.89</td>
</tr>
<tr>
<td>Mn (ppm)</td>
<td>463 ± 6</td>
<td>669 ± 11</td>
<td>520</td>
</tr>
<tr>
<td>Na (%)</td>
<td>3.06 ± 0.03</td>
<td>3.17 ± 0.03</td>
<td>3.17</td>
</tr>
</tbody>
</table>
Results

Five different geochemical types were identified among the 45 artifacts as shown in Figure 2a, Figure 2b and Table I. As shown in Figure 2a, the obsidian artifacts from Salmas were traced to two of the well-known Lake Van sources – one artifact came from the Nemrut source and six came from the Meydan source. There is little surprise here since the Lake Van sources are located closest to Salmas. However, as shown in Figure 2b, the artifacts from Kaleybar proved to be more interesting. Four different geochemical types were identified; however, only one of the artifacts could be traced to a known source. The single sample was traced to the Meydan obsidian source. The remaining 37 artifacts subdivide into three geochemical types: unknown #1 which has 31 artifacts, unknown #2 which has five artifacts, and unknown #3 which has one artifact. Comparisons of the NAA data from MURR with the data from Blackman et al. (1998) were unsuccessful in linking the artifacts to any of the known sources in Turkey, Armenia or Azerbaijan. Table II lists the calculated means and standard deviations for the three new obsidian source types identified during this investigation.

Discussion and Future Research

We conclude that the three newly established geochemical types are probably from sources in Iran (possibly Mt. Sahand or Mt. Sabalan) or from sources in the Transcaucasus that have yet to be reported. The Kaleybar site is relatively close to Mt. Sabalan and our data strongly suggest this area ought to be searched to locate possible obsidian deposits. The distance to Mt. Sahand is much greater but the area surrounding this volcanic peak should also be inspected to determine if any of the unknown artifacts at Kaleybar could have come from a source in that area.

Acknowledgements

The authors would like to acknowledge Mr. Bahman Kargar for providing the samples from Salmas. We also thank Dr. James Blackman who helped us by comparing the NAA data generated at MURR to his NAA obsidian database. The Archaeometry Laboratory at MURR is supported in part by the National Science Foundation (grant # 0504015). Any opinions, findings and conclusions or recommendations expressed in this material are those of the authors and do not necessarily reflect the views of the National Science Foundation.

References


A New Look at the Geochemistry of Obsidian from East Africa

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African Obsidian and the Origins of Modern Human Behavior

Databases of obsidian sources have been created for many parts of the world, but as of yet no comprehensive database exists for East Africa. Though some studies have been done on obsidian from this region, most notably Merrick and Brown (1984a,b & 1994), for the most part, they have not been comprehensive (Michels et. al. 1983; Negash et al. 2006a, 2006b, 2007). In addition, these data have not been collected in a manner conducive to the incorporation of new data collected using different analytical methods, and geographical coordinates for sources remain unpublished. However, these studies have shown the great potential for creating a comprehensive database for this region.

The work presented here is part of a continuing effort to create such a database by analyzing source samples from known sources to chemically characterize the geochemistry of the region, gathering the data from previous studies into one database to make it more easily accessible, and analyzing artifacts from this region to test the database and its ability to source artifacts accurately. The database will then be used to study obsidian source exploitation patterns to evaluate models of mobility, exchange, and interaction in ancient East Africa. This work focuses first on the analysis of the sources located in the Central Rift Valley in Kenya, a region rich in high-quality obsidian sources that were often used by ancient peoples.

One application of such a database is to examine the transition of humans to fully modern behavior (Ambrose 2001, 2002; Ambrose in press; Barut 1994). During the Middle Stone Age (MSA)/Late Stone Age (LSA) transition, we see a shift toward the development of socio-political organizations similar to ethnographically observed hunter-gatherers. This shift in organization is accompanied by changes in the patterns of raw material procurement. Typically, procurement of raw materials outside of the zone of direct procurement follows a distance-decay pattern, i.e. as the distance between a site and a source increases the relative abundance of that source decreases. However, as the interaction between socio-political groups increases, the slope of the typical distance-decay curve becomes shallower, as more artifacts from non-local materials are found at greater distances. Variation in mobility and home range size should also influence the slope of a curve and the size of the direct procurement zone. Hunter-gatherer home ranges are larger and social networks are more extensive and intensive in less productive, arid environments. Steeper slopes and smaller direct procurement zones should be found when higher rainfall supports more stable and productive environments (Ambrose 2002).

Figure 1. Location of obsidian sources in the central Rift Valley of Kenya analyzed in this study. Six chemical groups are plotted geographically. The white symbols represent chemically isolated samples.
Figure 2. A comparison between the XRF and NAA groupings. Figure 2A shows the results from XRF while Figure 2B shows the results from NAA. Both plots show the same groupings, though the groups formed with the NAA data are better resolved than those from the XRF data.

Methodology

The Central Rift Valley in Kenya was chosen as a good starting place, as there have been some studies done on obsidian from this general region, but there are many sources that are not sufficiently characterized (Ambrose in press). One hundred source samples were acquired by Dr. Stanley Ambrose of the University of Illinois for chemical analysis from 2001 to 2006 (Figure 1). The concentrations of trace elements are used to characterize the sources. The best elements are those incompatible with the solid phase due to large ion size or high ionic charges. One of the incompatible groups consists of the large-ion lithophile elements (LILE which include K, Rb, Cs, Sr, and Ba), the light rare earth elements (LREEs, La to Sm), and Th and U. Another of the incompatible groups consists of the high field-strength elements (HFSE which include Y, Zr, Hf, Nb, and Ta) and the heavy rare earth elements (HREEs, Eu to Lu). Many of these elements can be measured with excellent precision and accuracy using either XRF or NAA.

X-ray fluorescence (XRF) is a nondestructive technique that uses secondary x-rays to characterize the chemical composition of a sample. In XRF analysis, an x-ray of sufficient energy strikes the sample and is absorbed by the atoms causing the ejection of inner electrons. This puts the atom in an excited state and electrons from the outer shells are quickly transferred to fill the vacancies, giving off an x-ray in the process. These secondary x-rays are characteristic of the element that was struck by the primary x-ray and their energy is equal to the difference between the shell initially occupied by the electron and the shell it occupies after filling the vacancy. XRF is completely nondestructive, an important feature for the analysis of artifacts. The concentrations of eleven elements (K, Ti, Mn, Fe, Zn, Ga, Rb, Sr, Y, Zr, and Nb) in ppm are usually determined (Glascock 1998).

In neutron activation analysis (NAA), the sample is irradiated with neutrons, inducing radioactivity in the sample. These induced isotopes emit gamma rays as they decay to stable isotopes. The intensity of the gamma-ray emission is proportional to the original concentration of the element in the sample. NAA, due to the induced radioactivity, is a destructive technique. Typically, samples of about 100 milligrams are removed and used for analysis. Data acquired from NAA is typically more precise and more accurate than that obtained from XRF analysis. The relative concentrations of twenty-eight elements (Al, Ba, Ce, Co, Cs, Cl, Dy, Eu, Fe, Hf, K, La, Lu, Mn, Nd, Na, Rb, Sb, Sc, Sr, Sm, Ta, Tb, Th, U, Yb, Zn, and Zr) are determined by NAA (Glascock 1998). Neutron activation analysis was performed at the University of Missouri Research Reactor under the guidance of Dr. Michael Glascock.

For obsidian, simple bivariate plots can be used to identify source groups and compare compositional groups to geographic locations. The groups can then be validated statistically using Euclidean distances. Principal component analysis will also be used to
confirm that the best results are being acquired from the groups and the elements used to distinguish those groups, once there are enough samples to make the results statistically valid (Glascock 1998).

Figure 3. Plot of cesium versus iron. As can be seen most clearly in the group to the far right, some subgroups can be seen in some bivariate plots. However, more samples need to be analyzed to confirm the presence of these subgroups. In addition, there is a clear split in the iron concentration with the three groups on the left corresponding to the samples found south of the lake while the other groups correspond to the samples north of the lake.

Preliminary Results and Discussion

Thus far, the results are promising. The 100 source samples fall into at least six major chemically-distinct groups, with strong evidence for a few of those groups to be separated into subgroups. Plots of the rare earth elements and a few of the large-ion elements versus iron showed the greatest separation between groups. Looking at the XRF results, a plot of the log of the concentration of rubidium in parts per million (ppm) versus that of iron showed the best separation of the major groups (Figure 2). However, better resolution is needed in order to see subgroups. Using the NAA data, plots of cesium and the rare earth elements versus iron clearly show separation occurring within a few of the major groups, separating samples into subgroups.

In comparing these chemical source groups to their geographic locations, the samples fell into the same groupings with only a small amount of overlap between some possible subsources (Figure 1). Also, there appears to be a correlation between the concentration of iron and the location. Samples from the southern-most groups tend to have a lower concentration of iron (<3.0%) while northern samples tend to have a higher iron concentration (>4.5%) (Figures 1 and 3). In both cases, the iron concentration is relatively high, indicating that this region produced obsidian with peralkaline tendencies.

Future Work

Though the results of this study seem promising, a lot more work in this area must be done in order to distinctly characterize the geochemistry of this region of Africa as well as other regions. First, more samples will be collected from many of the sources examined in this study to confirm groupings, assess intra-source variation, and to improve statistics. Second, samples from outside the central Rift region will be collected to compare to these few original groups to gain a better understanding of the geochemistry of this area. Third, the results from previous studies done in this region will be compared with the results from this study to create a comprehensive database for these sources. Finally, artifacts will be obtained to test the completeness of the database and to begin applying the database to specific archaeological questions. Artifacts from several archaeological sites in Kenya do not match known sources (Merrick and Brown 1984; Merrick et al., 1994). An expanded source database would permit more comprehensive determination of site-to-source distances, and provide a firmer foundation for evaluating mobility, exchange and interaction patterns during the final stages of the evolution of modern human behavior. This database will be equally useful for understanding exchange and interaction patterns in later prehistory, particularly during the Neolithic era.
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Michels, J.W., I.S.T. Tsong, and C.M. Nelson

Negash, A., M. Alene, and M.S. Shackley


Negash, A., M. Alene, F.H. Brown, B.P. Nash & M.S. Shackley
Regional Scaling for Obsidian Hydration Temperature Correction

Alexander K. Rogers, Maturango Museum

Abstract

Hydration of obsidian is a temperature-dependent process, and correcting for temperature history is necessary for chronological use of obsidian hydration data. This paper describes a technique for estimating temperature history for an archaeological site, in which all the nearby sites in the same weather patterns are analyzed and a best scaling algorithm determined. The method uses data from publicly-available meteorological records. Two case studies are presented to illustrate the technique.

Introduction

It is well known that obsidian hydration is a temperature dependent process, and that temperature at an archaeological site varies both diurnally and annually (e.g. Friedman and Long 1976; Stevenson et al.1989, 1998, 2000, 2004; Rogers 2007). Thus, the effects of temperature history, manifested as effective hydration temperature (EHT) must be taken into account in any use of obsidian hydration data for chronological analyses. Computing a correction for temperature requires estimates of three temperature parameters: annual average temperature, annual temperature variation, and mean diurnal variation (Rogers 2007a). Furthermore, the temperature parameters must be representative of long-term conditions at the site; meteorological practice describes such data as norms, and typically demands a 30-year history (Cole 1970). Since most archaeological sites are not collocated with a modern weather station, estimating these parameters can present a problem.

This paper describes a technique of regional temperature scaling or estimation, in which all the nearby sites are analyzed and a best scaling algorithm determined. This has the advantage that it makes use of all the data available, and averages the effects of microclimates to some degree, with the caveat that the sites chosen must be in the same general weather patterns. The method is based on publicly-available meteorological records, from the web sites of the Regional Climate Centers. Two case studies to illustrate the technique are presented.

Alternative techniques involve temperature sensors at the archaeological site, or scaling from the nearest site, both of which have drawbacks. Temperature sensors placed at the site will yield short-term estimates, but these may not be representative of long-term conditions. Alternatively, estimates have been based on the nearest site with long-term temperature data, scaled for altitude by the mean adiabatic lapse rate (e.g. Gilreath and Hildebrandt 1997), but this procedure runs the risk that local microclimatic effects may affect the extrapolation. The method described herein avoids both these problems.

![Figure 1. Map of California, showing regions used for temperature scaling case studies.](image)
Analytical Technique

Computation of effective hydration temperature (EHT) is based on the equation (Rogers 2007a)

\[ EHT = T_a (1 - 3.8 \times 10^{-5} Y) + .0096Y^{0.95} \]  
(1a)
\[ Y = V_a^2 + V_d^2. \]  
(1b)

This requires three temperature parameters for the site: annual average temperature \((T_a)\); annual temperature variation \((V_a)\), defined as the difference between the July average temperature and the January average temperature; and mean diurnal variation \((V_d)\), defined as the average of the daily temperature ranges for July and January.

The two variation parameters, \(V_a\) and \(V_d\), represent the conditions to which an artifact was exposed, and thus are subject to corrections for depth (Carslaw and Jaeger, 1959; Rogers 2008; Stevenson et al. 1989) and for rock shelters (Everett-Curran et al. 1991; Rogers 2007a). The present analysis considers effects at the surface, so these corrections do not apply.

The validity of EHT as computed from equation 1 and this temperature model has been verified by comparison with EHT computed from actual hourly temperature data from the Amargosa Desert Research Site (Johnson et al. 2002, amplified by data through 2005); the standard deviation of the error between the two was \(<0.89^\circ C\) (Rogers 2008).

For the analysis here, temperature data for the desired sites are downloaded from the appropriate Regional Climate Center. The temperature parameters are then computed from the equations

\[ T_a = (\text{annual max temp.} + \text{annual min temp.})/2 \]  
(2)
\[ V_a = (\text{Jul max} + \text{Jul min})/2 - (\text{Jan max} + \text{Jan min})/2 \]  
(3)
\[ V_d = [(\text{Jul max} - \text{Jul min}) + (\text{Jan max} - \text{Jan min})]/2 \]  
(4)

and converted to °C. These parameters are then analyzed to determine the most appropriate and robust scaling method.

Two cases are described as examples; the locations are shown in Figure 1. The first is based on sites in the high desert of eastern California, and is typical of desert mountain regions with significant altitude relief. It spans a region roughly 200 km east-west by 300 km north-south, east of the Sierra Nevada, centered around a latitude of approximately 36°48’ N. The second is the central area of the Central Valley of California and its eastern foothills and Sierra slope, with Sacramento at its western edge, extending east to Nevada City. It extends roughly 200 km north-south and 100 km east-west, centered on a latitude of 38°30’ N. This region is typical of extended valleys and foothills, in which the climate is dominated by a temperature inversion layer. It will be seen that different scaling techniques are required for these two cases.

All the temperatures used in this study are air

<table>
<thead>
<tr>
<th>Station</th>
<th>Alt, ft</th>
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<th>Ave Min, deg F</th>
<th>Annual Ave, deg F</th>
<th>Jul Max, deg F</th>
<th>Jul Min, deg F</th>
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<td>73.6</td>
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<td>28.9</td>
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<td>101.9</td>
<td>66.0</td>
<td>60.2</td>
<td>31.7</td>
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<td>47.0</td>
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<td>100.6</td>
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<td>60.5</td>
<td>31.1</td>
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<td>63.4</td>
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<td>45.0</td>
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<td>92.3</td>
<td>65.3</td>
<td>51.1</td>
<td>29.4</td>
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<tr>
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<td>66.6</td>
<td>37.0</td>
<td>33.5</td>
<td>8.6</td>
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temperatures with 30 years of history, measured two meters above the ground in an enclosure which shelters the sensor from direct sunlight, normal meteorological practice.

High Desert

This case demonstrates a situation in which two of the three temperature parameters show strong scaling with site altitude. The analysis is based on monthly temperature data from the Western Regional Climate Center (WRCC), using the data base from 1971 – 2000. Table 1 summarizes the sites used in the temperature scaling analysis. All are from desert or desert mountain environments. The sites are all in the Mojave Desert/Great Basin weather patterns.

The data of Table 1 were used to compute the temperature parameters per equations 2 – 4; Table 2 summarizes the resulting parameters. Upon plotting, Figure 2 shows the scaling of $T_a$ with altitude, in which the data show a high degree of correlation with the straight-line fit. Note that the slope is $-1.8^{\circ}C/1000$ feet, which agrees well with the mean adiabatic lapse rate of $-1.9^{\circ}C/1000$ feet. Figure 3 shows the corresponding plot for $V_a$, again exhibiting a relatively tight grouping. (Generally differences between measured quantities are less numerically stable than averages because of random fluctuations; in this case the parameter is a difference of averages, and it appears to be fairly stable and predictable.) On the other hand, the plot of $V_d$ against altitude (Figure 4) shows very poor correlation. In this case the quantity $V_d$ again represents a difference of measured quantities, so the instability is not unexpected; in addition, the data represent short-term phenomena, which tend to be less stable than long-term data.

Figures 2 – 4 also display the linear best fit equation for each case. For $V_d$, the very small value of $R^2$ suggests that the best strategy is not to scale at all, but to use the mean value, $15.8^{\circ}C$, as the best estimate.

The data of Table 1 can also be used to examine microclimatic effects. Significant microclimatic differences exist between Inyokern and China Lake Armitage Field, which are within 15 km of each other and in the same valley. However, examination of the temperature parameter data in Table 2 suggests that such microclimates have little effect on $T_a$ and $V_a$, which both lie close to the best-fit curve (Figs. 2 and 3). The last parameter, $V_d$, does not track well between the two sites, but it is unstable anyway, as Figure 4 shows. Thus, if microclimatic effects are present, they will probably appear primarily in $V_d$, and even there it is not clear whether the differences are predictable.

Meteorological theory (e.g. Cole 1970) suggests that $T_a$ should scale for altitude by the mean adiabatic lapse rate, as shown in Figure 2. Figure 3 shows a similar scaling for $V_a$, suggesting that $T_a$ and $V_a$ should be related, so that knowing $T_a$ should allow predicting $V_a$. Figure 5 shows the strong correlation between $T_a$ and $V_a$, which demonstrates that $V_a$ can be predicted with considerable accuracy if $T_a$ is known.
Table 2. Temperature parameters calculated for high desert sites.

<table>
<thead>
<tr>
<th>Station</th>
<th>Alt, ft</th>
<th>Ta, deg C</th>
<th>Va, deg C</th>
<th>Vd, deg C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baker</td>
<td>940</td>
<td>21.17</td>
<td>21.56</td>
<td>17.67</td>
</tr>
<tr>
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<td>19.82</td>
<td>14.25</td>
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<tr>
<td>Daggett Apt</td>
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<td>20.61</td>
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<tr>
<td>White Mtn</td>
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<td></td>
<td></td>
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</tr>
</tbody>
</table>

Thus, for this desert case the annual average temperature was shown to be predicted by the equation

\[ T_a = 22.25 - 1.80x \]  

(5)

where \( x \) is altitude in thousands of feet. The error standard deviation of this model is 0.79°C, for the data set of Table 2. The annual temperature variation was found to decrease by 1.7°C/1000 ft. altitude increase, and to be predicted by

\[ V_a = 22.63 - 1.70x \]  

(6)

with \( x \) defined as above. The error standard deviation is 0.76°C.

As shown in Figure 4, the predictability of \( V_d \) with altitude is poor, so, in the absence of other data about a site, the most robust estimate is simply the mean:

\[ V_d = 15.8°C \]  

(7)

with an error standard deviation of 1.79°C. Finally, if \( T_a \) is known for a site, \( V_a \) is predicted by

\[ V_a = 1.65 + 0.94T_a \]  

(8)

with an error standard deviation of 0.27°C.

The purpose of the parameters is computation of effective hydration temperature (EHT) by equation 1. Figure 6 shows the EHT computed from equation 1 with equations 5 – 7 as input.
labeled “Best Fit”; it also shows EHT computed from equation 1 using the site meteorological data as inputs, labeled “Met data”. The error standard deviation of the between the two EHT estimates is 0.82°C. This accuracy is surprising because of the relatively large error in V_d; however, inspection of equation 1 shows that this arises because the EHT computation is dominated by the T_a value, which is characterized by smaller errors than V_d. The agreement between the two methods is remarkable.

**Figure 7.** Scaling of average annual temperature with altitude, Central Valley sites. Note break around 1500 ft.

**Central Valley**

The western slope of the Sierra Nevada presents a different case, in which less vertical relief exists and climate is dominated by a persistent temperature inversion layer in the Central Valley. The analysis is again based on monthly temperature data from the Western Regional Climate Center (WRCC), using the data base from 1971 – 2000. Table 3 summarizes the sites used in the temperature scaling analysis. All are in similar weather patterns. Temperature parameters were again computed by equations 2 – 4, and Table 4 summarizes the parameters.

**Figure 8.** Scaling of annual temperature variation with altitude, Central Valley sites.

Figure 7 shows the variation of T_a with altitude. As can be seen, the data for this case fall into two distinct patterns: valley floor and lower foothills, and higher foothills and mountains. The break point between the two is approximately 1500 feet altitude, probably correlating with the level of the normal inversion layer over the valley. Below this altitude, T_a is roughly independent of...
altitude, and has an average value of 15.85°C. Above 1500 ft the value of $T_a$ is given by

$$T_a = 19.44 - 2.7x, \quad 1500 < x < 4000 \quad (9)$$

where $x$ is altitude in thousands of feet.

Figure 8 shows the similar plot for $V_a$, again exhibiting a break at about 1500 ft. The average value of $V_a$ below 1500 ft is 16.02°C, while above 1500 ft the value is

$$V_a = 20.75 - 2.9x, \quad 1500 < x < 4000 \quad (10)$$

Finally, Figure 9 shows the scaling for $V_d$, with much poorer correlation with altitude, but noticeably better than the fit for the desert case. Analysis of accuracy shows that use of the altitude scaling gives better results than simply using the average, as was done for the desert. Thus, for sites below 1500 ft, the best fit is the mean value of $V_d$ is 14.77°C; above that altitude the best fit is

$$V_d = 18.57 - 1.5x, \quad 1500 < x < 4000 \quad (11)$$

Figure 7 and Figure 8 again show similar scaling with altitude for $T_a$ and $V_a$, and Figure 10 shows that there is again strong correlation between $T_a$ and $V_a$. However, there is a different grouping for valley floor and foothills, except this time the break point is below 100 ft. The equation relating $T_a$ and $V_a$ is

$$V_a = 1.21T_a - 3.57, \quad 0 < x < 100 \quad (12a)$$

Since the best fit to the data shows a break point, assessing parameter accuracy is more complex than for the desert case. Table 5 summarizes the accuracy achieved by this method, including a computation for effective hydration temperature made using equation 1. It can be seen that the algorithms yield values of $T_a$ and $V_a$ which are within 0.5°C, and $V_d$ within 1.7°C. Surprisingly, in view of the possible errors in $V_d$, the EHT values are within 0.5°C, again because the EHT computation is dominated by the $T_a$ value, which is characterized by the smallest errors.
### Table 4. Computed temperature parameters for Central Valley sites.

<table>
<thead>
<tr>
<th>Station</th>
<th>Alt, ft</th>
<th>Ta, deg C</th>
<th>Va, deg C</th>
<th>Vd, deg C</th>
</tr>
</thead>
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<td>Lodi</td>
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<td>Knights Ferry</td>
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<td>15.50</td>
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</table>

Figure 11 shows the values of EHT for the Central Valley sites, again computed from equation 1 with two different inputs. The solid line shows the values computed using the temperature model as summarized in Table 5, and the points show the EHT computed from the meteorological data for each site from Table 4. The agreement is obviously good.

The foregoing analysis suggests that for sites below approximately 1500 ft the best estimate of the temperature parameters is the mean for the site ensemble; above that altitude, altitude scaling should be employed.

### Discussions and Conclusions

The two case studies summarized above show that the form of the algorithm providing the best fit can only be determined from examination of the data, in conjunction with the physics of weather patterns, and may be different in different climatic regions. Judgment is required in doing this, and it is not possible to predict in advance what the best form of the temperature algorithms will be.

The method which gave the best results for high desert conditions was not the same as the best for the Central Valley conditions. In the desert, $T_a$ and $V_a$ scaled very accurately with altitude, while $V_d$ was best estimated by a mean value independent of altitude. In the Central Valley, none of the parameters for the sites below 1500 ft scaled with altitude, although altitude scaling was effective from 1500 – 4000 ft. A second difference was that, for the Central Valley sites, mean diurnal variation above 1500 ft has a positive correlation with altitude, so that use of altitude scaling yielded better accuracy in EHT than use of the mean.

The rates of decline of $T_a$ and $V_a$ with altitude varied as well between the two cases. In the desert, both approximated the mean adiabatic lapse rate, while in the Central Valley the rate of decrease with altitude above 1500 ft is much larger (-2.7°C/1000 ft for $T_a$ and -2.9°C/1000 ft for $V_a$). This more rapid decrease may be due to a sharp decline in humidity above the inversion layer, a phenomenon which does not occur in the desert.

### Table 5. Analysis of accuracy of temperature parameter models.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Site Altitude, ft.</th>
<th>Algorithm for Estimate, deg C</th>
<th>Mean Error, deg C</th>
<th>Error StDev, deg C</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T_a$</td>
<td>&lt; 1500</td>
<td>15.85</td>
<td>0.00</td>
<td>0.42</td>
</tr>
<tr>
<td></td>
<td>&gt; 1500</td>
<td>$T_a = 19.44 - 0.0027x$</td>
<td>0.02</td>
<td>0.14</td>
</tr>
<tr>
<td></td>
<td>Overall</td>
<td>-</td>
<td>0.01</td>
<td>0.32</td>
</tr>
<tr>
<td>$V_a$</td>
<td>&lt; 1500</td>
<td>16.02</td>
<td>0.00</td>
<td>0.34</td>
</tr>
<tr>
<td></td>
<td>&gt; 1500</td>
<td>$V_a = 20.75 - 0.0029x$</td>
<td>-0.06</td>
<td>0.23</td>
</tr>
<tr>
<td></td>
<td>Overall</td>
<td>-</td>
<td>-0.03</td>
<td>0.29</td>
</tr>
<tr>
<td>$V_d$</td>
<td>&lt; 1500</td>
<td>14.77</td>
<td>0.00</td>
<td>1.67</td>
</tr>
<tr>
<td></td>
<td>&gt; 1500</td>
<td>$V_d = 18.57 - 0.0015x$</td>
<td>0.11</td>
<td>0.95</td>
</tr>
<tr>
<td></td>
<td>Overall</td>
<td>-</td>
<td>0.04</td>
<td>1.42</td>
</tr>
<tr>
<td>EHT</td>
<td>&lt; 1500</td>
<td>Equation 1a,b</td>
<td>0.02</td>
<td>0.40</td>
</tr>
<tr>
<td></td>
<td>&gt; 1500</td>
<td>Equation 1a,b</td>
<td>0.04</td>
<td>0.06</td>
</tr>
<tr>
<td></td>
<td>Overall</td>
<td>Equation 1a,b</td>
<td>0.02</td>
<td>0.29</td>
</tr>
</tbody>
</table>
A final observation involves limits of validity of the resulting algorithms. First, the effects of paleoclimatic change have not been included; an implicit assumption was made that the parameters representing current temperature conditions are also valid through time. It is known that climatic shifts have occurred (West et al. 2007), but the effects of such shifts on obsidian hydration are generally small, < 7% in age (Rogers 2007b). A method for accounting for them has been described elsewhere (Rogers 2007b). Further, the algorithms for each case study were developed based on regions which are limited in geographical extent, in altitude, and in extent of general weather patterns. The algorithms should not be applied outside those limits without further analyses to verify validity - extrapolation is always dangerous.

In summary, regional analysis of temperature, based on publicly-available meteorological data, can yield valid algorithms for estimating temperature parameters at an archaeological site in the region. If an archaeological site is located in close proximity to a current weather station, the 30-year climate data from the station can be used to calculate the temperature parameters; otherwise, a regional temperature analysis should be performed as exemplified by the case studies in this paper. The resulting temperature parameters are representative of long-term present weather conditions, and provide the basis for accurate computations of effective hydration temperature in support of chronological analyses.

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