Time course of pharmacokinetic and hormonal effects of inhaled high-dose salvinorin A in humans

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Abstract
Salvinorin A is a kappa opioid agonist and the principal psychoactive constituent of the Salvia divinorum plant, which has been used for hallucinogenic effects. Previous research on salvinorin A pharmacokinetics likely underestimated plasma levels typically resulting from the doses administered due to inefficient vaporization and not collecting samples during peak drug effects. Six healthy adults inhaled a single high dose of vaporized salvinorin A (n = 4, 21 mcg/kg; n = 2, 18 mcg/kg). Participant- and monitor-rated effects were assessed every 2 min for 60 min post-inhalation. Blood samples were collected at 13 time points up to 90 min post-inhalation. Drug levels peaked at 2 min and then rapidly decreased. Drug levels were significantly, positively correlated with participant and monitor drug effect ratings. Significant elevations in prolactin were observed beginning 5 min post-inhalation and peaking at 15 min post-inhalation. Cortisol showed inconsistent increases across participants. Hormonal responses were not well correlated with drug levels. This is the first study to demonstrate a direct relationship between changes in plasma levels of salvinorin A and drug effects in humans. The results confirm the efficacy of an inhalation technique for salvinorin A.

Keywords
Salvia divinorum, salvinorin A, pharmacokinetics, prolactin, cortisol, endocrine

Introduction
The plant Salvia divinorum (a member of the mint family) has been used historically in shamanic practices of the Mazatec people of Oaxaca, Mexico for at least several hundred years (Ott, 1995; Valdés et al., 1983), although it was not botanically described until the 1960s (Epling and Jativa, 1962). Within the past 15 years S. divinorum has gained increased popularity as a psychoactive drug in non-traditional contexts (Perron et al., 2012; Wu et al., 2011). In non-traditional use, products containing S. divinorum leaves, sometimes infused with S. divinorum extract in order to increase drug effects, are typically smoked (Baggot et al., 2010; Gonzalez et al., 2006). Salvinorin A, the primary psychoactive compound in S. divinorum, is a kappa opioid agonist hallucinogen that is not active at the 5-HT2A receptor, the primary site of activity for classic hallucinogens such as lysergic acid diethylamide (LSD) and psilocybin (Cunningham et al., 2011; Prisinzano, 2005; Roth et al., 2002). Although S. divinorum and salvinorin A have not been controlled at the federal level in the US, at the time of this writing at least 35 states within the US and 27 nations have enacted various levels of restriction for S. divinorum (Siebert, 2015).

Understanding the effects of salvinorin A, including its pharmacokinetic profile, in humans is important for understanding recreational use of S. divinorum. Laboratory research has not found evidence of persisting psychotic-type episodes resulting from salvinorin A (Addy, 2012; Johnson et al., 2011; MacLean et al., 2013; Ranganathan et al., 2012). Cases of persisting psychotic-type episodes have been reported in association with recreational use, although the causal role of S. divinorum remains unclear (Vandrely et al., 2013). In addition, the dissociative and perceptual effects resulting from salvinorin A (Addy, 2012; Johnson et al., 2011; MacLean et al., 2013; Ranganathan et al., 2012) could potentially result in dangerous behavior in an unsupervised environment. Therefore, studying the pharmacokinetic profile of salvinorin A may inform the understanding of potential adverse reactions observed in recreational S. divinorum use.

Examining human salvinorin A effects is also important because salvinorin A or derivative compounds may serve as therapeutic agents for neurological (e.g. Alzheimer’s disease), pain, mood, personality, gastrointestinal, and cocaine-use disorders.
agonist administration (Ur et al., 1997). In order to address one aspect of the efficiency of the delivery system, residual salvinorin A from the glass pipe was assayed for each session.

Methods

Participants

Participants were 6 individuals who participated in a previous study assessing the effects of inhaled salvinorin A in the laboratory (Johnson et al., 2011; MacLean et al., 2013). The sample size was judged sufficient for examining pharmacokinetic data because robust significant subject-rated effects were observed with fewer participants (Johnson et al., 2011). Participants had taken part in up to 20 previous sessions (16 salvinorin A doses in ascending order and 4 intermixed placebo sessions under blind conditions) that did not involve collecting blood samples. Two individuals (one female and one male) whose subjective and cognitive data were included in our previous sample of eight participants (MacLean et al., 2013) did not participate in the final salvinorin A administration session, which was the only session involving blood draws. In the case of the male, the participant decided not to participate in the blood draw session upon considering several subjectively intense sessions previously in the study. In the case of the female, the investigators decided not to continue her onto the blood draw session due to excessive spontaneous arm movements in previous sessions, which may have interfered with the blood draws.

For the six participants reported here, the mean age was 25 years (range: 21–35). They reported using S. divinorum on a mean of 11 previous occasions (range: 1–40), with their reported first use at a mean age of 21 years (range: 16–31). They reported using classic hallucinogens on a mean of 32 previous occasions (range: 5–111). Study staff who were present during drug administration had established a rapport with participants during previous preparatory sessions and lower dose and placebo sessions as described previously (Johnson et al., 2011).

Procedure

Each participant inhaled a single high dose of vaporized salvinorin A. The dose administered was the highest tolerated dose of salvinorin A in previous sessions. For four participants, this dose was 21.0 mcg/kg, which was the maximal dose in the dose run-up. For the other two, this dose was 18.0 mcg/kg because they replied “yes” to a question asking them if they would refuse to receive the same or higher doses at the conclusion of a 19.5 mcg/kg session. As described previously (MacLean et al., 2013) subjective drug strength and monitor-rated effects (drug strength, distance from usual daily reality, unresponsiveness, psychological distress, paranoia, anxiety/fear, motor activity, joy/peace, and physical distress) and physiology measures (systolic and diastolic blood pressure and heart rate) were assessed every 2 min for 60 min after inhalation. Blood samples were collected at 13 time points (baseline, 1, 2, 3, 4, 5, 10, 15, 20, 25, 30, 60, and 90) and cold centrifuged to obtain plasma. Plasma samples were purified by solid phase extraction and analyzed in triplicate via liquid chromatography-tandem mass spectrometry (LC-MS/MS) using a +5 mass analogue of salvinorin A as the internal standard (Caspers et al., 2013). Analyses of prolactin and cortisol were performed with enzyme-linked immunosorbent assay (ELISA) kits (Calbiotech, Spring Value, CA) and were run in triplicate. The average of these three assays was used in the analyses. Residual salvinorin A in the glass pipe was determined for each session. Specifically, dichloromethane (1 mL in three separate washes) was used to wash the inner surfaces of the glass pipe. The combined 3 mL of resulting solution was then dried under a stream of nitrogen. Three separate samples from the resulting residue were dissolved into mobile phase and subjected to LC-MS/MS for analysis.
Data analysis

For each participant, we calculated Pearson’s correlations between each hormonal assay (prolactin and cortisol) and subjective and monitor ratings of drug effects, using only the 7 time points when both blood and ratings were collected (baseline, 2, 4, 10, 20, 30, and 60 min post-inhalation). Correlations were also conducted for each participant between salvinorin A and prolactin levels, between salvinorin A and cortisol levels, and between prolactin and cortisol levels. Correlations between drug and hormones used the 7 common time points indicated above, while correlations among hormones used 13 common time points. Because a delayed hormonal response (relative to drug levels) might obscure a relationship between drug and hormonal levels, the same pairs of correlations were also conducted at the group level using peak values (i.e. single maximal value across the time course for each participant) for drug and hormonal levels.

Repeated measures regression (SAS PROC MIXED, AR(1) covariance structure) was used to model the relationship between salvinorin A plasma level and participant and monitor ratings of drug effects from baseline to 60 min post-administration. As with the correlations, this analysis only used the 7 time points common to both blood draws and drug strength ratings. Statistical significance was defined as $p < .05$.

The percent of the intended dose that remained as residual salvinorin A in the glass pipe was calculated using the salvinorin A residual mass for each participant (i.e. mean of the triplicate LC-MS/MS assays) and the prepared absolute salvinorin A dose for each participant (i.e. taking bodyweight into account).

Results

Samples were collected and assayed for salvinorin A level at all time points. For prolactin and cortisol, sample volumes were insufficient to obtain results for two time points for one participant (at the 1 and 10 min time points). Coefficients of variation (CV) for plasma samples (in triplicate) and standards (in duplicate) were $< 10\%$. Figure 1(a) shows mean salvinorin A levels at all blood collection time points (up to 90 min post-inhalation). In order to show individual variability contributing to mean levels, Figure 1(b) shows individual participant salvinorin A levels at each time point up to 30 min post-inhalation. Figure 1(a) shows that mean peak salvinorin A levels occurred at 2 min post-inhalation, followed by rapid reductions and then more gradual reductions until the final time point at 90 min post-infusion, at which time salvinorin A levels were close to baseline (zero). Although these trends were generally observed at the individual participant level (Figure 1(b)), notable variations occurred, with peak effects occurring as early as 1 min to as late as 4 min post-inhalation.

To illustrate the relationship between drug blood levels and subjective drug strength, Figure 2(a)–(f) shows an individual participant’s salvinorin A plasma levels and subjective drug strength. Ratings of drug strength were closely associated with plasma levels. The median Pearson correlation between plasma levels and drug strength across individuals was $r = .93$ (range: .88–.99; all significant).

Repeated measures regression showed that salvinorin A level significantly increased participant $(F(1, 35) = 74.08, p < .0001)$ and monitor $(F(1, 35) = 29.14, p < .0001)$ ratings of drug strength and monitor ratings of distance from usual daily reality $(F(1, 35) = 15.41, p < .0001)$, unresponsiveness $(F(1, 35) = 19.82, p < .0001)$, psychological distress $(F(1, 35) = 21.26, p < .0001)$, and paranoia $(F(1, 35) = 11.87, p = .002)$. The effect of salvinorin A level was not significant for the remaining monitor ratings (anxiety/fear, motor activity, joy/peace, and physical distress). The effect of salvinorin A level was also not significant for physiology measures (systolic and diastolic blood pressure and heart rate). Results remained unchanged after controlling for lifetime use of hallucinogens and lifetime use of $S$. divinorum.

Figure 3 shows the effects of salvinorin A administration on plasma prolactin. Figure 3(a) shows mean prolactin levels, and Figure 3(b) shows prolactin levels in individual participants. Mean peak effects occurred at 15 min post-inhalation and gradually decreased through 90 min. However, individual participant data show a plateau of peak prolactin levels from 10 to 30 min post-inhalation for some individuals. Figure 4 shows the effects of salvinorin A administration on plasma cortisol. Figure 4(a) shows mean cortisol levels, and Figure 4(b) shows cortisol levels in individual participants. The mean cortisol time course resembled that of prolactin. However, there was substantial individual variability with little evidence of a cortisol response observed in some participants. There were no significant correlations between

![Figure 1](image-url)
Figure 2. Individual participant salvinorin A plasma levels (left axis) and subjective rating of drug strength (right axis) for all time points in which both measures were assessed. Individual participants are designated by the same symbols shown in Figure 1. The pre-inhalation assessment time point is shown at 0 min.
cortisol or prolactin levels and drug effect ratings within individual participants. No individual participant correlations between hormone levels and physiological measures were significant with the exception of one positive correlation between pulse and cortisol and one negative correlation between systolic blood pressure and prolactin. Salvinorin A levels were not significantly correlated with either cortisol or prolactin levels within any individual participant. Levels of prolactin and cortisol were positively correlated within each participant across the 13 time points (1 participant with 11 time points due to missing data) (Pearson r range: .36–.92; significant for 4 of 6 participants). In correlations at the group level, no significant relation was detected between salvinorin A and prolactin levels ($p = .82$), between salvinorin A and cortisol levels ($p = .15$), or between prolactin and cortisol levels ($p = .68$).

CV for the triplicates of residual salvinorin A assays for each participant were $< 4\%$. The mean mass of salvinorin A residue in the glass pipe across participants was 57.1 mcg (standard deviation (SD) $= 24.3$ mcg), representing a mean of 4.21% (SD = 2.25%) of the prepared absolute dose.

**Discussion**

This study is unique in that it examined the time course (including frequent, early time points) of salvinorin A plasma levels after salvinorin A inhalation, delivered via a relatively efficient vaporization system. The present study resulted in novel information relevant to three domains: drug delivery, time course of drug levels, and time course and magnitude of hormonal effects.

**Drug delivery**

The present study showed substantially higher salvinorin A plasma levels compared to the previous study of inhaled salvinorin A pharmacokinetics (Ranganathan et al., 2012). The previous study found a mean salvinorin A level of approximately 0.9 to 1.0 ng/mL resulting from 8 and 12 mg salvinorin A (with little difference between those two doses). In contrast, in the present study, at doses approximately eight times lower (18.0 and 21.0 mcg/kg, which equate to ~1.26 and 1.47 mg for a 70 kg bodyweight person), resulted in a mean of 18.8 ng/mL at peak effects. These data suggest the present
study used a substantially more efficient delivery method. Differences in efficiency could involve multiple factors including temperature and air flow topography. Moreover, the analysis showing only a small percentage of residual salvinorin A in the glass pipe highlights the efficiency of the delivery system.

**Time course of salvinorin A blood levels**

The present study found strong correspondence between salvinorin A levels and ratings of drug strength throughout the time course. Unlike the previous study of salvinorin A pharmacokinetics (Ranganathan et al., 2012), this study was able to demonstrate this relationship due to more frequent drug effect rating assessments and blood draws. The present results indicate that subjective effects of salvinorin A are a direct function of concurrent plasma levels of the drug. This finding is consistent with a study of intravenous salvinorin A in rhesus monkeys reporting overt sedation-like behavior effects generally overlapping with the period of detected plasma levels of salvinorin A (e.g. within ~15 min post-injection) (Schmidt et al., 2005).

**Time course and magnitude of hormonal response**

Similar to Ranganathan et al. (2012), the present study showed increases in prolactin and, less consistently, cortisol following salvinorin A administration. Due to infrequent sampling, the previous study did not have the ability to determine how closely hormone levels and salvinorin A levels were related in time. By showing rapid increases in salvinorin A levels that match the rapid subjective effects of the drug, the present study had the potential to demonstrate a strong correspondence between drug and hormone levels. However, the present study showed that prolactin and cortisol responses to salvinorin A administration followed a more delayed and prolonged time course than the drug itself.

**Conclusion**

This study provides important information regarding the pharmacokinetics of a relatively novel drug used for its hallucinogenic effects. It confirmed that a relatively efficient vaporization method resulted in substantially higher drug plasma levels compared to a previous study of salvinorin A pharmacokinetics (Ranganathan et al., 2012). Moreover, this study showed strong correlations between salvinorin A blood levels and drug strength ratings across the time course of drug effects, suggesting that subjective effects are a product of concurrent blood levels. This study also showed that salvinorin A generally increased prolactin, although it followed a more delayed and prolonged time course than the drug itself. Cortisol showed inconsistent increases across participants. Because smoking and vaporization both involve inhalation, the results of this study may be relevant to the recent use of *S. divinorum* in non-traditional contexts.

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