Acute toxicity of diphacinone in Northern bobwhite: Effects on survival and blood clotting

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Abstract

The anticoagulant rodenticide diphacinone was slightly toxic (acute oral LD₅₀ 2014 mg/kg) to Northern bobwhite (Colinus virginianus) in a 14-day acute toxicity trial. Precise and sensitive assays of blood clotting (prothrombin time, Russell’s Viper venom time, and thrombin clotting time) were adapted for use in quail, and this combination of assays is recommended to measure the effects of anticoagulant rodenticides. A single oral sublethal dose of diphacinone (434 mg/kg body weight) prolonged clotting time at 48 h post-dose compared to controls. At 783 mg/kg (approximate LD₉₀), clotting time was prolonged at both 24 and 48 h post-dose. Prolongation of in vitro clotting time reflects impaired coagulation complex activity, and was detected before overt signs of toxicity were apparent at the greatest dosages (2868 and 3666 mg/kg) in the acute toxicity trial. These clotting time assays and toxicity data will assist in the development of a pharmacodynamic model to predict toxicity, and also facilitate rodenticide hazard and risk assessments in avian species.

1. Introduction

Public concern over poisoning of non-target wildlife began in the early 20th century when strychnine and thallium used for rodent and predator control were observed to be affecting species like quail and song birds (Peterle, 1991). In the 1930s, Link investigated the hemorrhagic effects of improperly cured sweet clover (Melilotus spp.) in domestic livestock, and eventually isolated 3,3'-methylene-bis(4-hydroxycoumarin) (i.e., dicumarol) (Pelfrene, 1991). This discovery ultimately led to the synthesis of over 100 analogs, including warfarin that was eventually used as a rodenticide. The mechanism of action of warfarin, and other anticoagulant rodenticides, involves inhibition of synthesis of functional vitamin K-dependent factors including prothrombin.

Several current use anticoagulant rodenticides pose hazard to ground feeding birds by direct ingestion of bait (primary poisoning) and to predatory and scavenging birds by consumption of poisoned rodents or exposed invertebrate prey (secondary poisoning). Non-target exposure and effects of anticoagulant rodenticides in wildlife have been documented in many countries, including the United States, United Kingdom, France, Canada, Malaysia and New Zealand (e.g., Albert et al., 2009; Eason and Spurr, 1995; Howald et al., 1999; Lambert et al., 2007; Stone et al., 1999, 2003; Walker et al., 2008). For example, in New York State there were at least 51 confirmed cases of death by hemorrhage with detection of rodenticides in tissues of wildlife between 1971 and 1997, with over half of the incidents involving birds of prey (e.g., great horned owl, Bubo virginianus; red-tailed hawk, Buteo jamaicensis) (Stone et al., 1999). Based on these findings, an anticoagulant surveillance program was implemented between 1998 and 2001 for routine avian submissions to the New York State Wildlife Pathology Unit. Remarkably, 49% of 265 raptors had detectable quantities of anticoagulants in liver tissue, and anticoagulants were considered the cause of death in nearly 15% of these cases (Stone et al., 2003). On a global scale, the magnitude of the hazard of rodenticides to birds is not well-characterized because the effect of sublethal exposure on long-term survival and fitness is unknown, and most lethal poisoning events are probably unnoticed or not reported.

In 2002, the US Environmental Protection Agency (US EPA) completed a risk assessment that identified several rodenticides that posed a significant risk to birds and non-target mammals (Erickson and Urban, 2004). Subsequently, the US EPA (2008) released its final ecological risk mitigation decision that placed...
some restrictions on the sale, distribution, and packaging of several second generation rodenticides (i.e., brodifacoum, difethialone, bromadiolone and difencam) that have greater toxicity and are more persistent in tissues than first generation rodenticides. This action will likely be offset by expanded use of other anticoagulant rodenticides, including diphacinone (classified as a first or intermediate generation compound). Unfortunately, the hazard posed by diphacinone to non-target organisms is inadequately characterized. Median lethal dosages, sublethal responses and tissue concentrations are being generated in both traditional wildlife test species (e.g., Northern bobwhite, Colinus virginianus) and in birds of prey that are occasionally used in traditional wildlife toxicology programs (e.g., American kestrels, Falco sparverius) and Eastern screech owls, Megascops asio.

Fig. 1. Blood coagulation pathway in birds (adapted from Gentry, 1993 and Thomson et al., 2002: F=Factor, PL=phospholipid).

An acute oral toxicity study was conducted in bobwhite (weight range: 149–224 g) gavaged with technical grade diphacinone (99% active ingredient; Hacco, Inc. Randolph, WI) suspended in pure vegetable oil (soybean oil cholesterol-free; Crisco®, Orrville, OH). The selection of doses was somewhat problematic as existing data suggested that the slope of the dose–response curve was steep, and doses were too widely separated to permit estimation of a reliable median lethal dose (Campbell et al., 1991; US EPA, 1998). Accordingly, the study was conducted over a 6 week period using analytically determined diphacinone doses that were selected through an iterative process (917, 965, 1033, 2065, 2868 and 3666 mg/kg body weight; n=9–10 bobwhite/dose; approximately equal distribution of sexes/dose). Because of low solubility, diphacinone was administered as slurry in vegetable oil (heaviest in a dose group received 1 ml and the remaining birds received a fractional volume related to their weight, 0.76–0.99 ml). Some target doses could only be achieved by repeatedly administering 50 mg of target dose over a 24-h period (target exposure of 2868 mg/kg: 2 doses/day; target exposure of 2868 mg/kg: 3 doses/day and target exposure of 3666 mg/kg: 4 doses/day). Moderate volumes of the vehicle were also administered 1–3 times to controls (n=9) during a 24-h period. Birds were observed twice daily for signs of toxicity over a 14-day period.

Based upon the results of this acute toxicity trial, a second study was conducted in which bobwhite (weight range: 178–241 g) were gavaged with either vegetable oil (control n=6), or with analytically determined doses of diphacinone at 434 mg/kg (n=16) or 783 mg/kg (n=16). At 6, 12, 24 and 48 h post-dose, each bird was euthanized with carbon dioxide and immediately bled by cardiac puncture. Controls were sacrificed 48 h post-dose. Birds were sacrificed at 24 h post-dose by cervical dislocation. Blood samples (0.5 ml) were collected into syringes containing 50 μl 0.5 M EDTA, a suitable alternative to sodium citrate (Cérón et al., 2008). Blood samples were centrifuged, plasma harvested and frozen at −80 °C, and subsequently shipped to the PWRC for analysis.

2.2. Analytical determination of diphacinone in dosing solutions

For determination of diphacinone concentration in dosing solutions, 100 μl of the slurry were weighed and dissolved in 10 ml of 1:1 acetone:chloroform, and 10 μl was diluted to 10 ml with 5 mM tetrabutylammonium phosphate in 12 mM phosphate buffer (pH 8.5) and methanol (40:60 ratio). The sample was placed in an ultrasonic bath for 5 min, filtered through a 0.45 μm teflon syringe filter into an amber vial, and analyzed by reverse phase ion-paired high-performance liquid chromatography with concentrations determined by comparison to a calibration standard (Primus et al., 1996). The method detection limit was 4 ng/ml and the precision (mean coefficient of variation ± standard deviation) for triplicate determinations was 3.3 ± 1.5% (n=3).

2.3. Clotting assays

One-stage prothrombin time, Russell’s Viper venom time, and thrombin clotting time were measured using plasma samples from a third generation controlled BBL fibrometer (Becton Dickson & Co., Baltimore, MD) (Miale, 1965). Upon addition of reagents and plasma into a sample cup, a clot detection arm with a stationary electrode and a moving electrode automatically drop into the cup, and the formation of a fibrin clot between the electrodes (measurement endpoint) is determined by a detection circuit linked to a timer.

2.3.1. One-stage prothrombin time assay

In this assay, an excess of tissue factor and phospholipid (thromboplastin) interacts with plasma Factor VII to form an active complex, and through a cascade of subsequent reactions (Factor X activation, conversion of prothrombin to thrombin) fibrinogen is eventually converted to fibrin which forms the clot (Fig. 1). Crude chicken hatchling thromboplastin was prepared by the method of Quick as modified by Griminger et al. (1970) and Doerr et al. (1975). The cerebral cortex of 3-day old broiler chicks (Gallus gallus) was removed, weighed and homogenized in 10 volumes of cold acetone. This suspension was centrifuged at 1500g for 10 min, and the supernatant was discarded. The pellet was re-suspended in 5 ml of acetone, homogenized, centrifuged, acetone decanted, and this procedure was repeated. The pellet was then re-suspended in acetone, vacuum filtered using a Büchner funnel with a glass microfilter chamber (Whatman 934-AH, 1.5 μm retention, Fisher Scientific, Suwanee, GA). The material retained on the filter was dried over night in a vacuum desiccator, removed from the microfilter chamber, and ground individually in indoor pens (61 × 46 × 33 cm high) at NWRC. Bobwhite quail were maintained in a 12 h light: 12 h dark photoperiod at 18–21 °C, and provided food (Purina Game Bird Maintenance Chow® Product 5440, 12.5% protein, Denver, CO) and water ad libitum.
incubated with Simplastin\textregistered (rabbit brain thromboplastin; Trinity Biotech, Berkeley Heights, NJ). Intra-assay precision (mean coefficient of variation ± standard deviation) for duplicate determinations of bobwhite plasma with CHT was 4.5 ± 4.3% (n = 30). Inter-assay precision over the course of a year for human reference samples using Simplastin\textregistered was 2.9 ± 2.2% (n = 13). When the CHT solution was diluted, clotting time of bobwhite plasma was found to increase (Fig. 2). When bobwhite plasma was diluted with 8.3 mM sodium potassium phosphate buffer (pH 7.2), clotting time was relatively stable at plasma dilutions ranging between 40% and 80%, but increased dramatically at dilutions containing less than 20% plasma (Fig. 3).

2.3.2. Russell’s Viper venom time (RVVT)

In this assay, Russell’s Viper venom (RVV) directly activates Factor X (but not Factor VII) in the common pathway of the clotting cascade (Fig. 1). Reconstituted RVV Factor X activator (American Diagnostica, Stamford, CT) was diluted 1:10 with imidazole buffered saline (IBS; 0.0125 M imidazole 0.109 M NaCl, pH 7.4) and maintained at room temperature. Plasma (100 l) was incubated at 37 °C in a sample cup for 2 min, and 100 l of diluted RVV was added to the sample cup and incubated for 15 s. The clotting reaction was initiated with 100 l 25 mM CaCl\textsubscript{2}, and time to clot was measured (Tripplett and Harms, 1981a). Intra-assay precision for duplicate determinations of bobwhite plasma was 6.5 ± 13.5% (n = 19). Clotting time remained relatively stable when bobwhite plasma was diluted by as much as 60% with phosphate buffer, but RVVT increased dramatically at greater dilutions (Fig. 4).

2.3.3. Thrombin clotting time (TCT)

This assay measures the time for conversion of fibrinogen to fibrin (Fig. 1) using a standard thrombin solution (Tripplett and Harms, 1981b). This assay is insensitive to deficiency of vitamin K-dependent clotting factors but is an indicator of the amount of fibrinogen in the plasma sample. Fibrinogen deficiency from improper sample collection and handling can prolong prothrombin time and RVVT, and thus affect test method accuracy (increase false positive rate). For this heterologous assay, bovine thrombin reagent and human fibrinogen reference material was used (AMAX Fibrinogen kit, Trinity Biotech). A fibrinogen standard curve was prepared (65–520 mg/dL), and bobwhite plasma samples were diluted 1:10 with IBS. Diluted plasma (200 l) was incubated at 37 °C in a sample cup for 2 min, and the reaction initiated by the addition of 100 l of thrombin reagent. Clotting time of the test sample is transformed to fibrinogen concentration (mg/dL) from the fibrinogen standard curve. Intra-assay precision for duplicate determinations of bobwhite plasma was 5.7 ± 6.8% (n = 32).

2.4. Statistical methods

For the acute toxicity trial, the relation between diphacinone dose and survival, and estimates of the slope, median lethal dose (LD\textsubscript{50}) and 95% confidence interval were determined using the probit analysis procedure in SAS (SAS Institute, Carey, NC). In the sublethal dosing study, prothrombin time, RVVT and TCT measurement endpoints were tested for homogeneity of variance (Fmax test) and normality (Shapiro–Wilks test, normal probability plot and descriptive statistics). Homogeneity of variance and normality were improved following log-transformation, and measurement endpoints were then compared by analysis of variance (ANOVA) and a Tukey’s HSD test. A regression ANOVA was used to examine the relation between prothrombin time and dilution of the CHT. A one-phase decay model (Prism, GraphPad Software, Inc., La Jolla, CA) was used to describe the relation between clotting time and dilution of bobwhite plasma in the prothrombin time and RVVT assays.

3. Results

3.1. Acute and sublethal toxicity studies

Survival of bobwhite was significantly related (p < 0.0001) to dose of diphacinone (9 of 10 survivors at 917 mg/kg, 8 of 9 at 965 mg/kg, 10 of 10 at 1033 mg/kg, 7 of 10 at 2065 mg/kg, 1 of 10 at 2868 mg/kg and 0 of 10 at 3666 mg/kg), with most of the birds receiving greatest doses (2868 and 3666 mg/kg) dying within 1–3 days of exposure. Some of the dosed bobwhite seemed lethargic, and had fluffed feathers. Several of the birds that succumbed had subcutaneous bruises in the breast and back regions, which could reflect coagulopathy; however, there was no evidence of frank internal or external bleeding. Controls that had been dosed multiple times with vehicle in a 24 h period did not exhibit overt signs of toxicity. The LD\textsubscript{50} was estimated to be 2014 mg/kg (95% confidence interval 1620–2475 mg/kg), and the slope of the dose–response curve was steep (probit/log\textsubscript{10} ± standard error = 9.92 ± 2.27).
Based on this dose–response curve, the 783 mg/kg dose in the sublethal trial falls in the LD₀₁–LD₀₂ range, and the 434 mg/kg dose is just slightly greater than the lower 95% confidence limit of the LD₀₁. All bobwhite receiving these doses survived and did not exhibit overt signs of toxicity.

3.2. Clotting time

Administration of 434 and 783 mg diphacinone/kg body weight prolonged clotting time when compared to controls, but the temporal response was highly variable (Figs. 5 and 6). However, the fibrinogen concentration in 5 samples was undetectable, and the volume was too small to determine fibrinogen concentration in another sample. The absence of fibrinogen in these 5 samples suggests that they may have been collected improperly (viz., cardiac puncture of euthanized birds, blood sample partially clotted). Fibrinogen is generally in great excess, and its cleavage to fibrin in the TCT assay is not influenced by vitamin K antagonists. When these 6 samples were excluded, all remaining samples (n=32) contained more than 60 mg fibrinogen/dL plasma (range: 63–254 mg/dL). For these remaining samples, prothrombin time (Fig. 7) and RVVT (Fig. 8) were prolonged (p<0.05; factors of 3.9 and 4.4) at 48 h after administration of 434 mg/kg when compared to the control group. Following administration of diphacinone at 783 mg/kg, prothrombin time and RVVT were prolonged at both 24 h (p<0.05; factors of 2.5 and 2.8) and 48 h (p<0.05, factors of 4.1 and 3.6) when compared to the control group.

4. Discussion

Using categories of toxicity that describe harm (Loomis, 1978), results of the 14-day acute toxicity test suggest that diphacinone may be classified as only slightly toxic (LD₅₀ 2014 mg/kg) to Northern bobwhite. In a previous study (Campbell et al., 1991), a reliable median lethal dose for diphacinone in bobwhite could not be estimated (dosing concentrations were separated by a factor of 5), although inspection of the data suggest that the theoretical value fell between 400 and 2000 mg/kg (US EPA, 1998). Our estimated LD₅₀ in bobwhite is within the same order of
magnitude as that reported in mallards (Anas platyrhynchos; 3158 mg/kg) (Erikson and Urban, 2004). Based upon these data, technical grade diphacinone appears to be considerably less toxic to birds than to laboratory rats (range of estimated LD$_{50}$ 1.9–7 mg/kg), domesticated mammals (0.8–15 mg/kg), and wild mammals (0.2–340 mg/kg) (reviewed in Eisemann and Swift, 2006; Erikson and Urban, 2004). Accordingly, diphacinone baits or environmental residues would seemingly present a reduced hazard to birds compared to mammals. Nonetheless, it has been linked to secondary poisoning in raptors (Stone et al., 1999, 2003)

Furthermore, results of a recent 7-day acute oral toxicity study in American kestrels (F. sparverius) found diphacinone to be moderately toxic (LD$_{50}$ 97 mg/kg) (Rattner et al., 2010), suggesting that extrapolation of diphacinone toxicity from Northern bobwhite and mallards to other avian orders (e.g., Falconiformes, Stringiformes) may be dubious, and protection of raptors may require substantial safety factors.

Three coagulation assays were adapted that yielded short and precise clotting times with EDTA-treated plasma from bobwhite. Using a thromboplastin extract from chicken hatchlings, prothrombin time of untreated bobwhite (mean ± standard deviation: 15.2 ± 1.5 s, Fig. 7) was in the range of values reported for several species of domesticated birds (chicken; turkey, Meleagris gallopavo; Japanese quail, Coturnix coturnix) and wild birds (e.g., black ducks, Anas superciliosa; American kestrels; Eastern screech owls, M.asio; black kite, Milvus migrans; Egyptian vulture, Neophron percnopterus; wedge-tailed eagles, Aquila audax; pigeon, Columba livia; bronzing pigeons, Phaps chelcokera; Hispaniolan parrot, Amazona ventralis; umbrella cockatoo, Cacatua alba; Port Lincoln parrots, Barnardiacezonarius; Australian magpies, Gymnorhina tibicen) (Martin et al., 1994; Morrissey et al., 2003; reviewed in Powers, 2000; Rattner et al., 2009; Thomson et al., 2002; Webster, 2009). Using RVV that activates Factor X in the common pathway, clotting time of plasma from untreated bobwhite (39.5 ± 8.9 s, Fig. 8) was greater than that reported in several avian species (~9–21 s; Tahira et al. 1977; Timms, 1977), although precision for duplicate determinations (coefficient of variation 6.5%) and the standard deviation of the 5 control samples with >60 mg fibrinogen/mL seems acceptable. The volume of all plasma samples analyzed in the one-stage prothrombin time and RVVT assays was 100 μl, although dilution curves (Figs. 3 and 4) suggest that reagent incubation conditions permit much smaller volumes with only minimal effects on clotting time. Similar findings have been reported for prothrombin time assays in chicken (Bailey et al., 2005) and Japanese quail (Webster, 2009), and by using dilution curves, one could normalize clotting time of samples of variable volumes to a constant volume (e.g., 100 μl) (Bailey et al., 2005).

Anticoagulant rodenticides inhibit vitamin K-dependent post-translational processing of clotting Factors II (prothrombin), VII, IX and X (reviewed in Powers, 2000), but do not affect the synthesis of fibrinogen. Fibrinogen deficiency resulting from pathophysiologic conditions (e.g., hepatic synthetic failure, disseminated intravascular coagulation, fluid resuscitation after massive blood loss) or from improper sample collection practices can prolong in vitro clotting time. To date, avian studies examining anticoagulant rodenticide toxicity have failed to determine if sample fibrinogen content supports in vitro clot formation. Such a fibrinogen concentration threshold is yet to be established for birds, and in the interim, we used 60 mg/dL. Thus, a conservative diagnostic approach for anticoagulant rodenticide laboratory and field studies that evaluate vitamin K-dependent coagulopathies would entail a combination of assays, namely prothrombin time and/or RVVT, plus determination of fibrinogen to rule out any nonspecific influence on clotting time. Shlosberg and Booth (2006) have proposed that prolongation of prothrombin time by more than 25% is suggestive of anticoagulant exposure. Consideration should be given to standardizing avian thromboplastin preparations (e.g., International Normalized Ratios) and developing reference values for wild birds to facilitate interpretation of field exposure monitoring. In the absence of standardized avian thromboplastin preparations, RVVT offers the advantage of inter-laboratory comparisons as reagents are commercially available. Ideally, exposure should be confirmed by clotting time endpoints, fibrinogen concentration, and analytical detection of anticoagulant residues in blood or tissue.

In the present study, several samples did not contain detectable quantities of fibrinogen, failed to clot in both the one-stage prothrombin time and RVVT assays, and thus, were excluded from our evaluation of the effects of diphacinone on hemostasis. Clotting time of samples with detectable fibrinogen (>60 mg/dL) was not affected at 6 and 12 h after administration of diphacinone, but was prolonged at 24 and 48 h post-dosage. This time course corresponds with, and may precede, the onset of overt toxicity and lethality at high dosage levels used in the acute toxicity study. The lag time between dosing and development of coagulopathy reflects the half-life clearance of functional coagulation factors and the increasing circulation of des–carboxy, dysfunctional factors. Diphacinone’s irreversible inhibition of hepatic vitamin K epoxide reductase impairs post-translational carboxylation of the prothrombin group of serine protease coagulation factors (Factors II, VII, IX and X). Without the addition of a carboxyl group, these factors are unable to assemble on cell surfaces to form active coagulation complexes. Related studies in American kestrel have shown a similar time course for prolonged clotting time following diphacinone administration (peak effect at 48 h post-dose), and also demonstrated that clotting time returns to control levels by day 7 post-exposure (Rattner et al., 2010). Detection of prolonged prothrombin time is used as an early indicator of anticoagulant rodenticide ingestion in domestic mammals, and prothrombin time is used routinely to monitor coumadin anticoagulant therapy in people (Mount and Feldman, 1983; Spinler et al., 2005). Prolonged prothrombin times have been reported within days of: (1) dietary exposure to diphacinone (American crow, Corvus brachyrhynchos, Massey et al., 1997; golden eagles, Aquila chrysaetos, Savarie et al., 1979), warfarin (chickens, Veltmann et al., 1981) and brodifacoum (Japanese quail, Webster, 2009), (2) repeated gavage with pindone (wedge-tailed eagles, bronzing pigeons, Port Lincoln parrots, black ducks, Australian magpies, Martin et al., 1994) and (3) intravenous injection of warfarin (chickens, Stopforth, 1970). Aside from evoking outright mortality due to hemorrhage, prolonged clotting time constitutes a change in fitness that could affect tolerance to normal physiological events (e.g., blood loss associated with molt) and environmental stressors (e.g., exposure to contaminants that impair hematopoiesis, altered nutritional plane).

5. Conclusion

Diphacinone was found to be of low acute oral toxicity to Northern bobwhite, with effects on coagulation detectable at sublethal concentrations. A series of clotting assays were developed and applied for use in bobwhite that are sensitive, precise, inexpensive, linked to the pathogenesis of toxicity (and ultimately mortality), and have applicability as biomarkers in both controlled laboratory studies and field monitoring. Determination of prothrombin time and/or Russell’s Viper venom time, plus fibrinogen concentration to exclude nonspecific influences on clotting, is recommended for diagnostic purposes. These findings and assay methods will assist in the development of a pharmacodynamic model to predict toxicity, and also facilitate rodenticide...
hazard and risk assessments of secondary poisoning in avian species.

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