Abstract: Llama derived single domain antibodies (sdAb), the recombinantly expressed variable heavy domains from the unique heavy-chain only antibodies of camels, were isolated from a library derived from llamas immunized with a commercial abrin toxoid preparation. Abrin is a potent toxin similar to ricin in structure, sequence and mechanism of action. The selected sdAb were evaluated for their ability to bind to commercial abrin as well as abrax (a recombinant abrin A-chain), purified abrin fractions, Abrus agglutinin (a protein related to abrin but with lower toxicity), ricin, and unrelated proteins. Isolated sdAb were also evaluated for their ability to refold after heat denaturation and ability to be used in sandwich assays as both capture and reporter elements. The best binders were specific for the Abrus agglutinin, showing minimal binding to purified abrin fractions or unrelated proteins. These binders had subst $\text{NM}$ affinities and regained most of their secondary structure after
heating to 95 °C. They functioned well in sandwich assays. Through gel analysis and the behavior of anti-abrin monoclonal antibodies, we determined that the commercial toxoid preparation used for the original immunizations contained a high percentage of Abrus agglutinin, explaining the selection of Abrus agglutinin binders. Used in conjunction with anti-abrin monoclonal and polyclonal antibodies, these reagents can fill a role to discriminate between the highly toxic abrin and the related, but much less toxic, Abrus agglutinin and distinguish between different crude preparations.

**Keywords:** abrin; single domain antibody; reversible refolding

**Figure S1.** Luminex direct binding of llama anti ricin (L<R) and llama anti abrin (L<A) polyclonal antibodies as well as phage displayed sdAb. Binding examined for commercial abrin (Abrin), abrax (Aac), ricin, ricin a chain (Rac) and ricin B chain (Rbc). Clones 1B, H11, and G12 were negative by monoclonal phage ELISA and are also negative by the Luminex direct binding assay.

**Figure S2.** Amino acid sequence determined from DNA sequencing of the unique anti-abrin sdAb characterized for this work. These 10 clones represent 8 sequence families.
**Figure S3.** Sandwich assays showed the Abr 2 and N12 bound to the same epitope on the *Abrus* agglutinin, but N5 bound to a different epitope than N12. Left panel: A ProteON GLC chip coated with *Abrus* agglutinin was exposed to the sdAb N12 at 30 nM in lanes 2–5 (light blue, dark blue, green and pink), lane 1 (red) and 6 (orange) were buffer blanks. Right panel: To evaluate epitope location the same area was challenged with Abr 2 30 nM (lanes 1–3) and N5 30 nM (lanes 4–6). Abr 2 generated a signal on lane 1 but not 2 or 3 indicating it recognized the same epitope as N12, while N5 generated a signal on all 3 lanes indicating it recognized a different epitope than Abr 2.

**Figure S4.** SPR data with fits for the binding of the selected sdAb to commercial abrin, abrax, *Abrus* agglutinin or abrin fractions. Time, in seconds, is on the X axis, while the Y axis if the SPR signal reported as resonance unit s (RU). Data and the calculated fits are shown. “No Binding” indicates that no binding was observed by SPR.
**Figure S5.** Sandwich assays for the detection of commercial abrin (left), mixture of the abrin fractions (middle), and *Abrus* agglutinin (right) using sdAb and mAb captures paired with the indicated sdAb tracer. Captures are as follows: Orange triangle N12; blue square mAb 5F6; red circle and green diamond llama anti abrin; blue diamond mAb 30x anti-ricin. Concentration, in ng/mL, is on the X axis and signal, reported as median fluorescence intensity (MFI), is on the Y axis.
Figure S6. Native (top left) and denaturing (top right and bottom) gels. Labeled markers are shown on the left of each gel. In the top panels lanes 1–3 are abrin fractions I, II, III; lane 4 is *Abrus* Agglutinin; lane 5 is the commercial abrin; lane 6 is ricin and Lane 7 is RCA120. The bottom panel shows isolated recombinant abrin A chain (Abrax), analyzed by SDS-PAGE and Coomassie staining. The purified protein is 29 kDa consistent with the coding sequence of the abrin A chain.