

Tracking *Neospora caninum* parasites using chimera monoclonal antibodies against its surface antigen-related sequences (rNcSRS2)

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1 **Tracking *Neospora caninum* Parasites Using Chimera**
2 **Monoclonal Antibodies against its Surface Antigen-related**
3 **Sequences (rNcSRS2)**

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14 (ABSTRACT)

15 Neosporosis, an infectious disease of cattle and dogs, causes an abortion in
16 cattle, which has a major damage on the dairy industry worldwide. Tracking of
17 *Neospora caninum* parasite that is responsible for neosporosis is required for the
18 prevention of this infectious disease. We developed a chimera monoclonal antibodies
19 consist of variable regions of murine antibody and constant regions of human
20 antibody against *N. caninum*. Recombinant surface antigen-related sequence 2
21 (rNcSRS2) of *N. caninum* was expressed in silkworm larvae, and immunized in mice
22 to obtain phage displaying antibody library. Through three rounds of selection,
23 three antibodies, A6, E1 and H3, were isolated and bound to rNcSRS2 with
24 nanomolar to micromolar affinity. In immunofluorescent staining assays, A6 and E1
25 bound to *N. caninum* strain Nc-Liv, demonstrating a successful tracking of the
26 parasite. H3 clone bound to rNcSRS2 but not to a truncated protein without
27 glycosylphosphatidylinositol (GPI) anchor domain in the carboxyl terminal. Amino
28 acid sequences of A6 and E1 were similar, but that of H3 differed in the CDR-H1
29 region, which might be the reason of their difference of affinity. These antibodies are
30 thought to be useful for prevention of cattle from neosporosis.

31 **Keywords:** neosporosis, parasite, antibody, phage display, immunofluorescent staining

(INTRODUCTION)

32

33 Neosporosis is a coccidian disease caused by *Neospora caninum* in a variety of
34 animals such as cattle, horses, deer, dogs, and so on (1), induces an abortion in cattle,
35 which is the reason for the huge economic damage on the dairy industry in many
36 countries (2, 3). The major mode of transfection of *N. caninum* is transplacental parasite
37 transmission that infects in the herd over successive generations (4, 5), but horizontal
38 transmission between cattle was not observed. There are three infectious stages in the life
39 cycle of *N. caninum*: tachyzoite, tissue cysts, and oocysts. Tachyzoites and tissue cysts
40 are the stages found in the intermediate hosts and they occur intracellularly (6); the
41 unsporulated oocysts are found in domestic dogs, which are the only known definitive
42 host for *N. caninum* (7).

43 Like other coccidian parasites, tachyzoites of *N. caninum* also contain the
44 characteristic organelles such as dense granules, rhoptries, and micronemes. Those
45 organelles produce important proteins that play important roles in infection of host cells
46 (8). On the other hand, proteins located on the surfaces are considered to play very
47 important roles in the infection process. They may induce the interaction with the host
48 cell and subsequently help the parasites adhere to and invade the host cell. Surface
49 antigen 1(NcSAG1) of *N. caninum* is the immunodominant surface antigen of tachyzoites,

67 of phagemid, and TG-1 for displaying antibody on M13 phage were purchased from
68 Agilent Technologies (La Jolla, CA, USA). Phagemid pDong1/Fab (16), helper phage
69 KM13 and non-suppressor *E. coli* strain HB2151 were provided kindly by Dr. Hiroshi
70 Ueda of Chemical Resources Laboratory of Tokyo Institute of Technology. The *N.*
71 *caninum* Nc-Liv strain (No. 50845) and Vero cell (No. CCL-81) were purchased from
72 ATCC (Rockville, MD, USA). Restriction and modification enzymes were purchased
73 from Takara-Bio (Shiga, Japan), Toyobo (Osaka, Japan), Roche Diagnostics (Tokyo,
74 Japan), or New England Biolabs (Tokyo, Japan). Oligonucleotides were synthesized
75 either by Operon (Tokyo, Japan) or Invitrogen (Tokyo, Japan). Other chemicals, reagents,
76 and antibodies, unless otherwise indicated, were obtained from Sigma-Aldrich (St Louis,
77 MO, USA) or Wako Pure Chemical (Osaka, Japan).

78 **Immunization of mice with rNcSRS2** The strategy for developing monoclonal
79 antibodies is shown in [Fig. 1A](#). Firstly, mice were immunized with rNcSRS2. After the
80 quantitation of peptide-specific antibodies in sera, the variable region genes of the
81 antibody heavy (V_H) and light (V_L) chains were prepared and cloned to a phagemid
82 vector to perform phage display selection. The rNcSRS2 was expressed in silkworm
83 according to previous report (15) and purified. Two inbred BALB/c mice (Japan SLC, Inc.
84 Hamamatsu, Shizuoka, Japan) were immunized with purified rNcSRS2 four times at

85 2-week intervals with a dose of 100 μ g through the subcutaneous route. The protein
86 solutions were emulsified with a Freund's complete adjuvant (Rockland Immunochem,
87 Gilbertsville, PA, USA) to increase the efficiency of immunization. After the last
88 immunization, blood samples were taken by tale bleeding and the rNcSRS2-specific
89 antibodies in sera were confirmed by an Enzyme-Linked ImmunoSorbent Assay (ELISA)
90 with immobilized rNcSRS2 on a microplate as described by Dong et al. (17). The
91 experiments with animals were carried out in the Animal House of Shizuoka University
92 in strict accordance with the recommendations in the Guide for the Care and Use of
93 Laboratory Animals of Shizuoka University and were approved by the Committee on the
94 Ethics of Animal Experiments of Shizuoka University (Permit Number: 24-11).

95 **Construction of phage display antibody library** The total RNA was
96 extracted from spleen cells of immunized mice with TRIzol (Invitrogen, Tokyo, Japan).
97 The genes of V_H and V_L of antibodies were amplified using PrimeScript One step
98 RT-PCR kit ver.2.0 (Takara, Shiga, Japan) according to the manufacturer's protocol. The
99 mouse IgG-specific primers are synthesized based on the common antibody primer
100 sequences (17). The PCR products were then purified using IllustraTM GFXTM PCR DNA
101 and Gel Band Purification kit (GE Healthcare). The purified V_L fragments were digested
102 with restriction enzymes *SalI* and *NotI* and were purified and ligated into a phagemid

103 pDong1/Fab digested with the same enzymes using T4 DNA ligase at 16°C for 1 h. After
104 confirmation of the inserted V_L sequence of several clones out of the obtained ones, the
105 V_H genes were inserted into the V_L-inserted phagemid library using restriction enzymes
106 *Sfi*I and *Xho*I. Electroporation-competent *E. coli* TG-1 cells were transformed with the
107 ligation product and plated on 2×YTAG agar (16 g/l tryptone, 10 g/l yeast extract, 5 g/l
108 NaCl, pH 7.2, supplemented with 100 µg/ml ampicillin, 1% glucose, and 1.5% agar)
109 plates overnight at 37°C. The size of library was estimated from the number of colonies
110 on the plate. *E. coli* TG-1 cells, transformed with the phagemid, were cultivated in 4 ml
111 of 2×YTAG overnight at 37°C. Ten milliliters of 2×YTAG were inoculated with 100 µl of
112 the overnight culture at 37°C at 200 rpm until OD₆₀₀ reached ~0.5, when helper phage
113 KM13 (18) was added with a multiplicity of infection (MOI) of 20. After incubation at
114 37°C for 30 min without shaking, the culture was centrifuged at 3700 g for 15 min. The *E.*
115 *coli* pellet was resuspended in 50 ml of 2×YTAK (2×YT medium containing 100 µg/ml
116 ampicillin and 50 µg/ml kanamycin) and incubated overnight with shaking at 30°C. The
117 overnight culture was centrifuged at 10,800 g for 30 min. Ten milliliters of PEG/NaCl
118 solution (20% polyethylene glycol 6000, 2.5 M NaCl) was added to 40 ml of supernatant,
119 and the mixture was incubated on ice for 1 h. After incubation, the mixture was
120 centrifuged at 6,000 g for 30 min. The pellet was resuspended in 2 ml of PBS and

121 centrifuged at 15,000 g for 10 min to pellet cell debris, and the supernatant was collected
122 as a Fab-displaying phage solution.

123 **Enrichment and selection of monoclonal antibodies** For antibody selection, 100
124 μl of rNcSRS2 (1 $\mu\text{g}/\text{ml}$ in PBS) was immobilized on a microplate at 4°C overnight. It
125 was washed three times with 200 μl of PBST, and then blocked with MPBS (PBS
126 containing 2% skim milk) for 2 h, followed by adding 10^{12} colony forming unit (cfu)
127 phage in 100 μl PBS and incubated for 1 h at room temperature. Phages bound to the
128 microplate were eluted with 100 μl of 1.0 mg/ml TPCK-treated trypsin (Sigma-Aldrich)
129 in PBS after washing with PBST for six times. *E. coli* TG-1 cells ($\text{OD}_{600} = 0.5$ in 700 μl
130 culture) were infected with 100 μl of eluted phage solution and cultured in 10 ml of
131 2×YTAG medium at 37°C with shaking at 200 rpm. When OD_{600} reached 0.5, the KM13
132 helper phage was added at an MOI of 20, and incubated for 30 min at 37°C without
133 shaking. After being centrifuged at 3,700 g for 10 min, the pellet was resuspended in 50
134 ml of 2×YTAK medium and incubated with vigorous shaking at 30°C overnight. The
135 culture supernatant was prepared by centrifugation at 10,800 g for 30 min, and phages
136 were precipitated with 0.2 volume of PEG/NaCl on ice for 1 h. After centrifugation at
137 6,000 g for 30 min, the pellet was resuspended in PBS and used as a source of Round 1
138 (R1) phage. Round 2 (R2) antibody selection from R1 phage was performed as described

139 above and R2 phage was obtained. From the R2 phage library, Round 3 (R3) selection
140 was also carried out to generate R3 phage. The enrichment of rNcSRS2-specific
141 phage-antibody among the original phage library (R0), R1, R2, and R3 phages was
142 confirmed with a polyclonal phage ELISA.

143 After the increase of binding capability of phage was confirmed, 96 infected *E. coli*
144 clones at the 3rd biopanning were picked up and cultivated for making monoclonal phage.
145 A phage ELISA was performed for 96 individual clones to select rNcSRS2-specific
146 phage-antibodies. Nucleotide sequence of positive clones was read by Greiner Bio One
147 (Tokyo, Japan) with primer M13rv (5'-GGAAACAGCTATGACCATG-3') for V_H and
148 primer VLseq (5'-CACTGGCTGGTTTCGCTAC-3') for V_L, and was analyzed using a
149 GENETYX software (Genetyx Corporation, Tokyo, Japan).

150 **Expression of Fab fragments of positive clones** pDong1/Fab was designed for the
151 convenient expression of the Fab fragment after selection of positive clone by placing a
152 TAG amber codon (19) between V_H-C_{H1} and gene III of phage. With this design, Fab is
153 expressed as a fusion protein with protein gIII of phage in suppressor strain like *E. coli*
154 TG1, resulting in the display of Fab on the surface of phage. However, with a
155 non-suppressor strain, such as *E. coli* HB2151, the Fab fragment will be expressed as a
156 soluble fragment (16). Because the genes for C_{H1} and C_L gene in pDong1/Fab system

157 were from human IgG (C_H1 and C_κ), Fabs expressed in this study were murine-human
158 chimera fragments. In brief, 200 μl of exponentially growing *E. coli* HB2151 was
159 infected with 10⁹ cfu of phage for 30 min at 37°C. Infected *E. coli* cells were pelleted by
160 centrifuge at 5,000 g for 10 min, resuspended in 4 ml 2×YT medium containing 100
161 μg/ml of ampicillin (2×YTA), and cultivated for 3 h at 37°C. Four hundred milliliters of
162 2×YTA medium was inoculated with the 4-ml culture and cultivated at 37°C with shaking.
163 Once the OD₆₀₀ reached 0.5, isopropyl β-D-thiogalactoside (IPTG), with a final
164 concentration of 1 mM, was added and cultivated further overnight at 30°C. The *E. coli*
165 cells were harvested by centrifugation at 4,000 g for 20 min at 4°C. The periplasmic
166 fraction was extracted according to a general protocol. His-tagged Fabs were purified
167 from the periplasmic fraction and concentrated supernatant with TALON
168 Co²⁺-immobilized resin (Takara-Bio) according to the instructions provided by the
169 manufacturer. Because one-step purification was not enough to achieve purity, Fabs were
170 furthermore purified with an anti-FLAG M2 affinity gel (Sigma-Aldrich) according to the
171 instructions provided by the manufacturer. The purified Fabs were analyzed using
172 SDS-polyacrylamide gel electrophoresis as described by Laemmli (20).

173 **ELISA analysis** The antigen-binding capacity of phage-displayed Fab fragments
174 was tested with ELISA. The microplates (NUNC, Langenselbold, Germany) were coated

175 overnight with 100 μ l of rNcSRS2 (0.5 μ g/ml) per well or 10 μ g/ml of BSA in PBS at
176 4°C. The plate was blocked at 25°C for 2 h with 2% MPBS, washed three times with
177 PBST, and incubated with 100 μ l/well of MPBS containing 10^9 – 10^{10} cfu of
178 Fab-displaying phage at 25°C for 1 h. The plate was washed three times with PBST and
179 incubated with 100 μ l/well of 5000-fold diluted HRP/anti-M13 monoclonal conjugate
180 (from Sheep; GE Healthcare UK limited, Little Chalfont, Buckinghamshire HP7 9NA,
181 UK) in MPBS at 25°C for 1 h. The plate was then washed three times with PBST and
182 developed with 100 μ l/well of 3,3',5,5'-tetramethylbenzidine (TMBZ) substrate solution.
183 After incubation for an appropriate time, the reaction was stopped by adding 50 μ l/well of
184 10% sulfuric acid, and the absorbance was read using a Model 680 microplate reader
185 (Bio-Rad, Hercules, CA, Japan) at 450 nm with 655 nm as a control.

186 For evaluation of IC₅₀ of phage Fabs, competitive ELISAs were performed in which
187 free rNcSRS2 in a series of concentration levels competed with immobilized rNcSRS2 to
188 bind Fab-phages.

189 For confirmation of antigen-binding capacity of free Fab fragments, a Rabbit
190 anti-Human Kappa chain antibody (1 μ g/ml; MBL, Tokyo, Japan) and an ECLTM
191 Anti-mouse IgG, Horseradish Peroxidase linked whole antibody (from Sheep; GE
192 Healthcare UK Limited) were used.

193 **Surface plasmon resonance (SPR) analysis** Binding analysis of three chimera
194 monoclonal antibodies to rNcSRS2 was performed using BIAcore X-100 or 2000 (GE
195 Healthcare Japan, Tokyo, Japan). In brief, purified recombinant NcSRS2 from silkworm
196 larvae (1800-4000 RU) was immobilized on the CM5 sensor chip (GE Healthcare Japan)
197 by amine-coupling method under pH 4.0. Serially diluted chimera monoclonal antibodies,
198 A1, E2 at concentrations of 78, 312, 625 and 1250 nM, H3 at 179, 358, 715, 1430 and
199 2860 nM, was injected to the sensor chip with HBS-EP buffer (10 mM HEPES, 150 mM
200 NaCl, 3 mM EDTA, 0.005% surfactant P-20 (GE Healthcare Japan), pH 7.4) at 30 μ l/min,
201 respectively. Ten mM Glycine buffer (pH2.5) was used as a regeneration buffer.

202 **Immunofluorescent staining of parasite with Fab fragments** *N. caninum* strain
203 Nc-Liv was used to infect Vero cells. The infected cells were maintained in a CO₂
204 incubator at 35°C. For immunofluorescent staining, freshly purified parasites were fixed
205 on glass slides coated with amino silane (APS), permeabilized, and blocked with 4%
206 BSA in PBS. Immunolabeling was carried out using 10 μ g/ml of purified Fabs. A
207 commercial anti-*N. caninum* antibody (1 μ g/ml; VMRD, Pullman, WA, USA) for positive
208 control was used. A sample without any primary antibody was used as a negative control.
209 After 1-h incubation, a rabbit anti-human kappa chain antibody (1 μ g/ml; MBL, Tokyo,
210 Japan) was added to samples on the glass slide, except positive control.

211 Rhodamine-conjugated goat IgG fraction anti-rabbit IgG (MP Biomedicals, LLC-Cappel
212 Products, Santa Ana, CA, USA) in 200 times dilution and 4',6-diamidino-2-phenylindole
213 (DAPI; 1000 times dilution; DOJINDO, Kumamoto, Japan) were added to all samples,
214 except for positive control after incubation, and the samples were washed with PBS. For
215 positive control, Alexa Fluor 594-conjugated AffiniPure Rabbit Anti-Mouse IgG (H+L)
216 (1 µg/ml; Jackson ImmunoResearch Laboratories, West Grove, PA, USA) and DAPI
217 were added. All samples were viewed on a confocal microscope (LSM 700, Carl-Zeiss,
218 Oberkochen, Germany), and their images were processed by employing the software
219 ZEN lite 2010 (Carl-Zeiss).

220

221

RESULTS

222 **Immunization of mice** Two inbred BALB/c mice were used for immunization.
223 After the last immunization, blood samples were taken to check the *N. caninum*
224 protein-specific antibodies. Specific binding capacity of sera was observed against
225 rNcSRS2, but not to BSA which was used as negative control, suggesting that the
226 immunization was successful (data not shown).

227 **Monoclonal antibody selection from phage display library** The display of the

228 Fab fragment on the surface of phage was achieved by using pDong1/Fab with the help of
229 the KM13 helper phage. For construction of antibody library, the V_H and V_L genes of
230 antibodies were amplified and detected at 350–400 bp in an agarose electrophoresis (data
231 not shown). A phage display antibody library with a diversity of 5×10^6 was obtained.
232 After three rounds of selection, the enrichment of rNcSRS2-binding phage was confirmed
233 using an ELISA with original phage library R0 and sublibraries R1, R2, and R3, which
234 were amplified in each step of biopanning. Absorbance at 450 nm in phage ELISA for R0,
235 R1, R2, and R3 phage against rNcSRS2 rose with the increase of the biopanning step (Fig.
236 1B), suggesting that three rounds of biopanning enriched the rNcSRS2-specific
237 Fab-phages. These phage pools did not bind to BSA as a negative control.

238 R3 phages were used to infect *E. coli* TG-1 for forming colonies. Ninety-six colonies
239 were picked up and cultivated for forming phage. Four clones A6, E1, H3, and H6
240 showed a strong signal against immobilized rNcSRS2 (Fig. 2A). The binding capacity of
241 those Fab-phages was also reconfirmed against rNcSRS2 and BSA. A6, E1, and H3
242 clones bound to rNcSRS2 but not to BSA (Fig. 2B), suggesting their specificity to our
243 target. Although H6 clone bound to rNcSRS2 in the screening, it did not bind to rNcSRS2
244 upon further confirmation (data not shown). The KM13 helper phage in which no
245 antibody was displayed was used as a negative control and did not bind to immobilized

246 rNcSRS2.

247 **IC₅₀ of A6, E1, and H3 clones** To evaluate the half maximal inhibitory
248 concentration (IC₅₀) of those clones, a competitive ELISA with serially diluted rNcSRS2
249 solutions inhibiting the binding of Fab-phage to immobilized rNcSRS2 was performed.
250 Competition was observed between free and immobilized proteins, and the IC₅₀ values of
251 A6, E1, and H3 were evaluated to be 0.54, 0.22, and 0.51 µg/ml of rNcSRS2,
252 respectively (data not shown).

253 **Purification of Fab antibodies and their binding to rNcSRS2** Fab antibodies
254 were expressed in *E. coli* HB2151 and purified with TALON Co²⁺-immobilized resin and
255 an anti-FLAG M2 affinity gel. The purified Fab fragments showed two bands on
256 SDS-PAGE with molecular weight of 24 and 26 kDa (Fig. 3A), which were identified as
257 light chain and V_H-C_{H1} of heavy chain, respectively. The binding of these Fabs to
258 rNcSRS2 and a truncate rNcSRS2 without GPI anchor domain of native protein rNcSRS2
259 (NSNT) (unpublished data) was confirmed with an ELISA. As shown in Fig. 3B, all the
260 three Fabs bound to the rNcSRS2 and A6, E1 also bound to rNcSRS2 (NSNT). However,
261 the H3 clone did not bind to the truncated protein, suggesting it might bind to the anchor
262 domain of the native protein. All Fabs did not bind to BSA as a negative control.

263 **Antigen binding affinity of monoclonal antibodies** Rate constants of
264 association (k_a) and dissociation (k_d), and dissociation constant (K_D) for each antibody
265 against the immobilized rNcSRS2 were investigated by SPR. Using the binding
266 sensorgrams (Fig. 4), k_a , k_d , and K_D are summarized in [Table 1](#). The K_D for clone H3 was
267 70 nM, much lower than A6 and E1 due to its low k_d . K_D values of A6 and E1 clones
268 were 420 nM and 5 μ M, respectively. The difference in absolute rate constants reflect
269 their different structure, especially that of complementarity determining regions which
270 determine the antibody's avidity and specificity for specific antigens.

271 **Immunofluorescence assay of *N. caninum* with monoclonal antibodies** To
272 confirm whether those clones bind to *N. caninum* parasites, immunofluorescence assays
273 were performed using a commercial anti-*Neospora* antibody as a positive control. As
274 shown in [Fig. 5](#), the parasites' nuclear was identified with DAPI-staining, showing blue
275 fluorescence under confocal laser microscopy. Staining with the A6 and E1 Fabs, a
276 Rabbit anti-Human Kappa chain IgG and Rhodamine-conjugated Goat IgG fractions
277 anti-Rabbit IgG revealed red fluorescence, suggesting both A6 and E1 Fabs bound to
278 parasites as well as the commercial anti-*N. caninum* antibody did. For the negative
279 control without primary antibody, no fluorescence was observed. In this experiment, H3
280 clone did not bind to parasites.

298 were also analyzed. The sequences of these antibodies were identified to be novel. Amino
299 acid sequences of A6 and E1 were similar; however, a change of one amino acid in the
300 CDR-H1 region caused different affinity between A6 and E1. H3 clone has a quite
301 different sequence with the A6 and E1 especially in CDR regions, which might bind to a
302 different epitope of rNcSRS2.

303 In immunofluorescent staining assays, both A6 and E1 bound to the tachyzoites of
304 the parasite. SRS protein of *Toxoplasma gondii*, sharing similar characteristics with *N.*
305 *caninum*, has been reported to be involved in the infection of parasite to host cells (21).
306 Therefore, there is a high possibility that these monoclonal antibodies can be used as a
307 blocker of infection. Monoclonal antibodies are now established as a key therapeutic
308 modality to a range of disease. Owing to the ability of these agents to selectively target
309 tumor cells, cancer has been a major focus of development programs for monoclonal
310 antibodies (22). Nowadays, many antibody drugs for humans have been developed and
311 have obtained approval from governments. However, there are few for animal therapy.
312 The antibodies developed in this study also provide a possibility for the development of
313 antibody drugs against neosporosis. Recently, antibody fragments such as Fab and
314 single-chain variable fragment (scFv) are more enthusiastically studied as drugs because
315 they have small molecular weights and bear the advantages of easy detection and tracking

316 in vivo, as well as the easy delivery to certain cells through antibody specificity. By
317 adding the Fc region gene and transforming insect cell or mammal cells, the whole length
318 antibodies for some certain applications can also be achieved.

319 Even though H3 clone bound to rNcSRS2, it did not bind to the truncated protein
320 without an anchor domain. rNcSRS2 was reported to be expressed in insect cell using
321 baculovirus expression system with all the necessary post-translational modifications (21),
322 however, the process of GPI anchor has not been revealed yet. Therefore, this clone may
323 be useful for studying the mechanism of the GPI anchor process.

324 Nowadays, most detection methods for neosporosis focus on the detection of
325 antibodies in the serum sample of cattle. However, approaches for monitoring the
326 infection of *N. caninum* in dairy farm could be achieved because antibodies developed in
327 this study can directly detect *N. caninum* parasites. This would further protect livestock
328 from infection of *N. caninum* parasite, leading to a development of antibody drugs for *N.*
329 *caninum* therapy.

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336 **References**

- 337 1. **Dubey J.P.:** Review of *Neospora caninum* and neosporosis in animals, Korean J.
338 Parasitol., **41**, 1-16 (2003).
- 339 2. **Cramer G., Kelton D., Duffield T.F., Hobson J.C., Lissemore K., Hietala S.K.,
340 and Peregrine A.S.:** *Neospora caninum* serostatus and culling of Holstein cattle, J.
341 Am. Vet. Med. Assoc., **221**,1165-1168 (2002).
- 342 3. **Dubey J.P.:** Recent advances in *Neospora* and neosporosis, Vet. Parasitol. **84**,
343 349-367 (1999).
- 344 4. **Davison H.C., Otter A., and Trees A.J.:** Estimation of vertical and horizontal
345 transmission parameters of *Neospora caninum* infections in dairy cattle, Int. J.
346 Parasitol., **29**,1683-1689 (1999).
- 347 5. **Schares G., Peters M., Wurm R., Barwald A., and Conraths F.J.:** The efficiency
348 of vertical transmission of *Neospora caninum* in dairy cattle analysed by serological
349 techniques, Vet. Parasitol., **80**, 87-98 (1998).
- 350 6. **Dubey J.P., Barr B.C., Barta J.R., Bjerkas I., Bjorkman C., Blagburn B.L.,
351 Bowman D.D., Buxton D., Ellis J.T., Gottstein B., Hemphill A., Hill D.E., Howe**

- 352 **D.K., Jenkins M.C., Kobayashi Y., Koudela B., Marsh A.E., Mattsson J.G.,**
353 **McAllister M.M., Modry D., Omata Y., Sibley L.D., Speer C.A., Trees A.J.,**
354 **Uggla A., Upton S.J., Williams D.J., and Lindsay D.S.:** Redescription of *Neospora*
355 *caninum* and its differentiation from related coccidia, *Int. J. Parasitol.*, **32**, 929-946
356 (2002).
- 357 7. **Lindsay D.S., Dubey J.P., and Duncan R.B.:** Confirmation that the dog is a
358 definitive host for *Neospora caninum*, *Vet. Parasitol.*, **82**,327-333 (1999).
- 359 8. **Dubey J.P., Carpenter J.L., Speer C.A., Topper M.J., and Uggla A.:** Newly
360 recognized fatal protozoan disease of dogs, *J. Am. Vet. Med. Assoc.*, **192**,1269-1285
361 (1988).
- 362 9. **Boothroyd J.C., Hehl A., Knoll L.J., and Manger I.D.:** The surface of
363 *Toxoplasma*: more and less, *Int. J. Parasitol.*, **28**, 3-9 (1998).
- 364 10. **Jung C., Lee C.Y., and Grigg M.E.:** The SRS superfamily of *Toxoplasma* surface
365 proteins, *Int. J. Parasitol.*, **34**, 285-296 (2004).
- 366 11. **Nishikawa Y., Xuan X., Nagasawa H., Igarashi I., Fujisaki K., Otsuka H., and**
367 **Mikami T.:** Monoclonal antibody inhibition of *Neospora caninum* tachyzoite
368 invasion into host cells, *Int. J. Parasitol.*, **30**, 51-58 (2000).
- 369 12. **Jacquet A., Coulon L., De Neve J., Daminet V., Haumont M., Garcia L., Bollen**

- 370 **A., Jurado M., and Biemans R.:** The surface antigen SAG3 mediates the
371 attachment of *Toxoplasma gondii* to cell-surface proteoglycans. *Molecular and*
372 *biochemical parasitology*, **116**, 35-44 (2001).
- 373 13. **Lekutis C., Ferguson D.J., Grigg M.E., Camps M., and Boothroyd J.C.:** Surface
374 antigens of *Toxoplasma gondii*: variations on a theme, *Int. J. Parasitol.*, **31**,
375 1285-1292 (2001).
- 376 14. **Nishikawa Y., Inoue N., Xuan X., Nagasawa H., Igarashi I., Fujisaki K., Otsuka**
377 **H., and Mikami T.:** Protective efficacy of vaccination by recombinant vaccinia
378 virus against *Neospora caninum* infection, *Vaccine* **19**, 1381-1390 (2001).
- 379 15. **Otsuki T., Dong J., Kato T., and Park E.Y.:** Expression, purification and
380 antigenicity of *Neospora caninum*-antigens using silkworm larvae targeting for
381 subunit vaccines, *Vet. Parasitol.*, **192**, 284-287 (2013).
- 382 16. **Dong J., Ihara M., and Ueda H.:** Antibody Fab display system that can perform
383 open-sandwich ELISA, *Anal. Biochem.*, **386**, 36-44 (2009).
- 384 17. **Dong J., Otsuki T., Kato T., Kohsaka T., Ike K., and Park E.Y.:** Development of
385 Two Murine Antibodies against *Neospora caninum* Using Phage Display Technology
386 and Application on the Detection of *N. caninum*, *PloS one*, **8**, e53264 (2013).
- 387 18. **Litman G.W., Rast J.P., Shablott M.J., Haire R.N., Hulst M., Roess W., Litman**

388 **R.T., Hinds-Frey K.R., Zilch A., and Amemiya C.T.:** Phylogenetic diversification
389 of immunoglobulin genes and the antibody repertoire, *Molecular biology and*
390 *evolution*, **10**, 60-72 (1993).

391 19. **Weigert M.G., Lanka E., and Garen A.:** Amino acid substitutions resulting from
392 suppression of nonsense mutations. II. Glutamine insertion by the Su-2 gene;
393 tyrosine insertion by the Su-3 gene, *J. Mol. Biol.*, **14**, 522-527 (1965).

394 20. **Laemmli U.K.:** Cleavage of structural proteins during the assembly of the head of
395 bacteriophage T4, *Nature*, **227**, 680-685 (1970).

396 21. **Nishikawa Y., Tragoolpua K., Makala L., Xuan X., and Nagasawa H.:** Neospora
397 caninum NcSRS2 is a transmembrane protein that contains a
398 glycosylphosphatidylinositol anchor in insect cells, *Vet. Parasitol.*, **109**, 191-201
399 (2002).

400 22. **Reichert J.M., and Valge-Archer V.E.:** Development trends for monoclonal
401 antibody cancer therapeutics, *Nat. Rev. Drug. Discov.*, **6**, 349-356 (2007).

402

403

404 **Figure Legends**

405 FIG. 1. Scheme for the development of murine anti-*N. caninum* monocloal antibodies

406 (A) and the enrichment of rNcSRS2-specific clones (B). ELISA was performed to
407 confirm the enrichment of rNcSRS2-specific clones. In ELISA, rNcSRS2 (0.5 µg/ml) and
408 BSA (10 µg/ml) were immobilized on a 96-well microplate, respectively. HRP/anti-M13
409 monoclonal antibody conjugate was used as the secondary antibody. R0 stands for the
410 original phage library, whereas R1, R2, and R3 stand for the amplified Fab-phage in
411 Rounds 1, 2, and 3 of biopanning, respectively.

412 FIG. 2. Screening of monoclonal antibodies (A) and their bindings to rNcSRS2 and
413 BSA. Protein rNcSRS2 (0.5 µg/ml) were immobilized on a 96-well microplate,
414 respectively. HRP/anti-M13 monoclonal antibody conjugate was used as the secondary
415 antibody after the addition of a 96-phage Fab solution (A). rNcSRS2 (0.5 µg/ml) and
416 BSA (10µg/ml) were immobilized on a 96-well microplate, respectively. HRP/anti-M13
417 monoclonal antibody conjugate was used as the secondary antibody after addition of
418 phage Fab solution (B). Experimental data were presented as average values with
419 standard error ($n=3$).

420 FIG. 3. SDS-polyacrylamide gel electrophoresis analysis of purified Fab antibodies
421 (A) and their bindings to rNcSRS2, rNcSRS2 (NSNT), and BSA (B). For ELISA,
422 rNcSRS2, rNcSRS2 (NSNT) (0.5 µg/ml each), and BSA (10 µg/ml) were immobilized on

423 a 96-well microplate, respectively. After incubation with Fab antibodies, a Rabbit
424 anti-Human Kappa chain antibody (1 µg/ml) and an ECLTM Anti-mouse IgG, Horseradish
425 Peroxidase linked whole antibody (from sheep) was used as secondary and tertiary
426 antibodies, respectively. M: Precision Plus ProteinTM Dual Colors Standards.

427 FIG. 4. SPR sensorgrams of the purified monoclonal antibodies, A6 (A), E1 (B) and
428 H3 (C), bound to recombinant NcSRS2. Recombinant NcSRS2 was immobilized on the
429 CM5 sensor chip and serially diluted antibodies were injected to the sensor chip.

430 FIG. 5. Immunofluorescence assay of *N. caninum* with monoclonal antibodies. *N.*
431 *caninum* strain Nc-Liv was immobilized and blocked followed by the addition of a
432 commercial anti-*N. caninum* antibody (P), Fabs A6, E1, and H3, respectively. A sample
433 without any primary antibody was a negative control (N). For Fab samples and negative
434 control, a Rabbit anti-Human Kappa chain antibody (1 µg/ml) was used as secondary
435 antibody and Rhodamine-conjugated Goat IgG fraction Anti-Rabbit IgG was added
436 finally. For positive control, Alexa fluor 594-labelled anti Mouse IgG was used as a
437 secondary antibody. 4',6-Diamidino-2-phenylindole (DAPI) was used to stain the nuclear
438 of parasites. The samples were observed under a confocal laser-scanning microscope.
439 Scale bar stands for 10 µm.

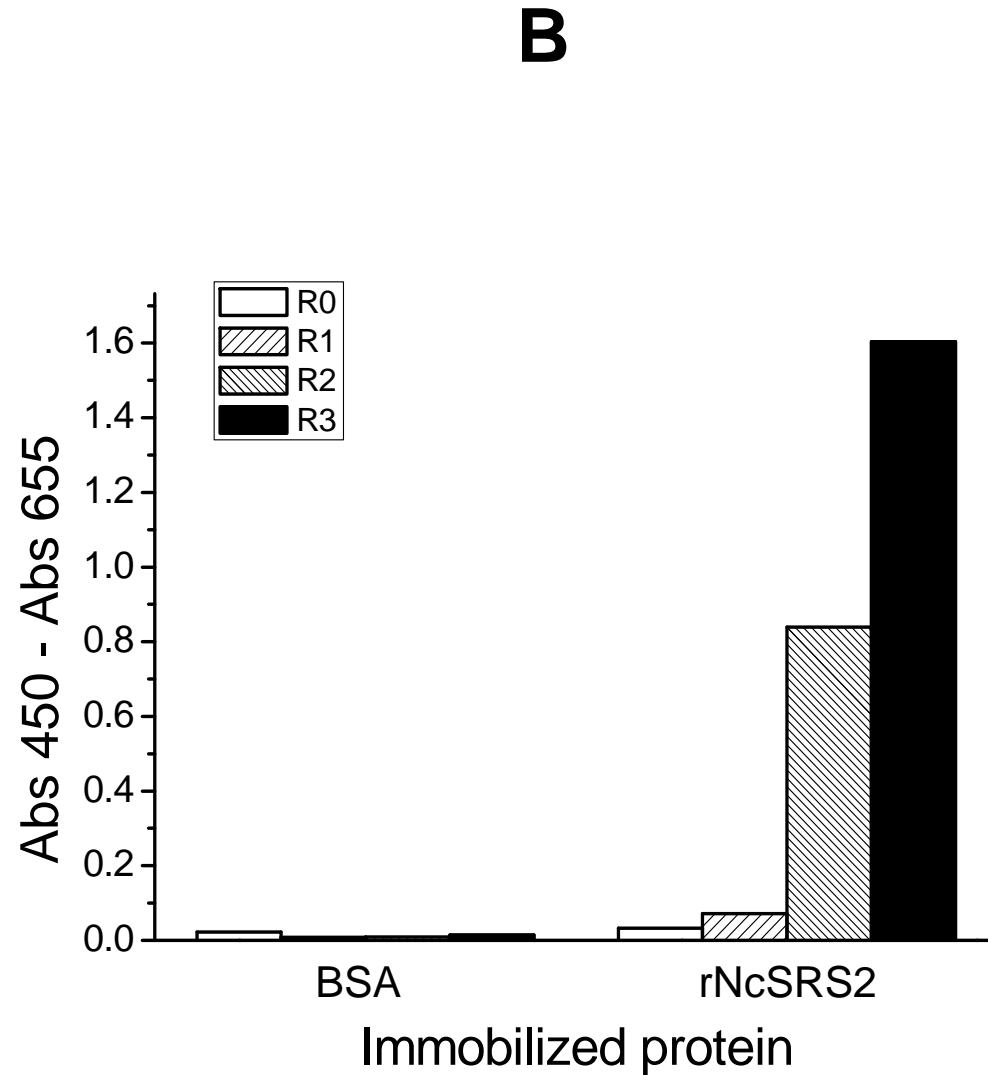
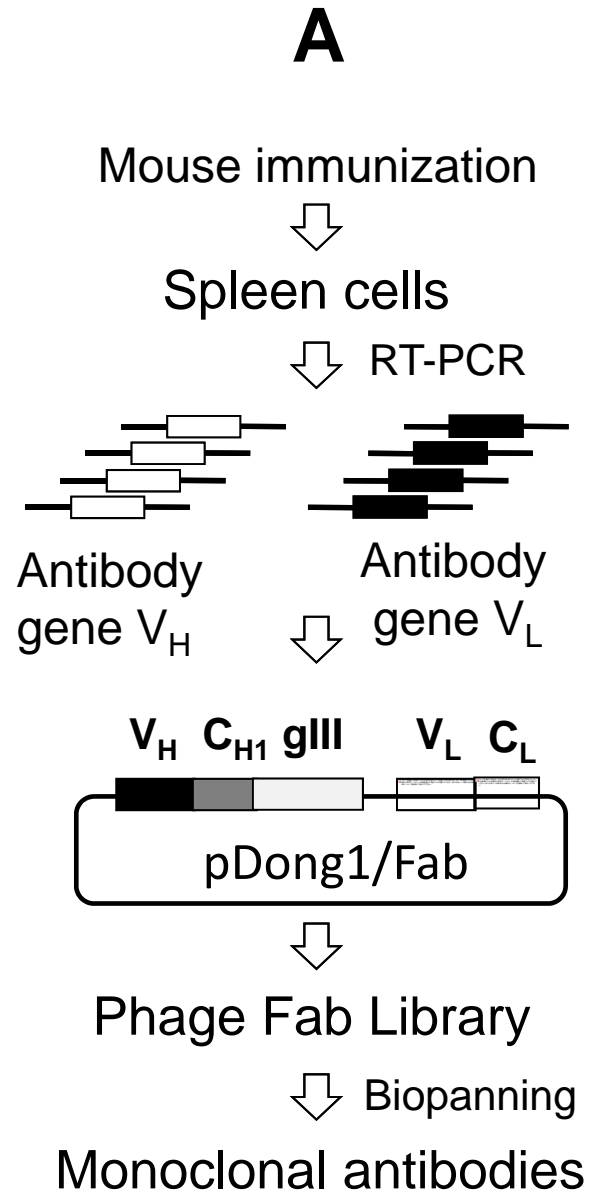
440 FIG. 6. Amino acid sequences of anti-NcSRS2 antibody variable regions. The amino
441 acid numbers are according to the Kabat numbering scheme. The amino acids of
442 complementarity determining regions (CDRs) are bolded and the different amino acids
443 between A6 and E1 are highlighted.

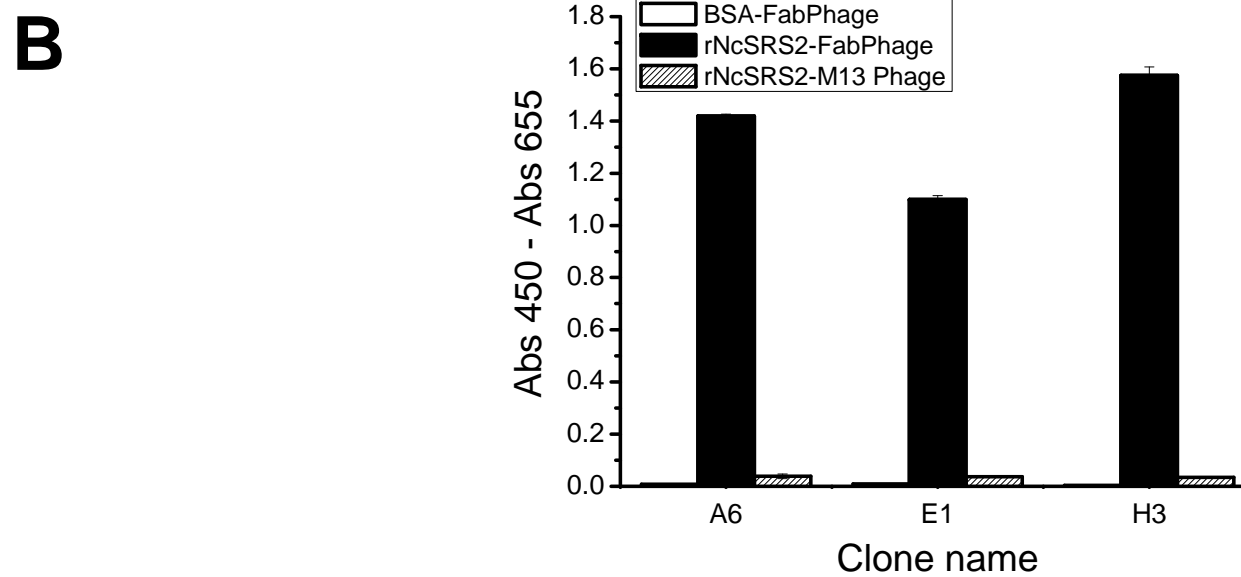
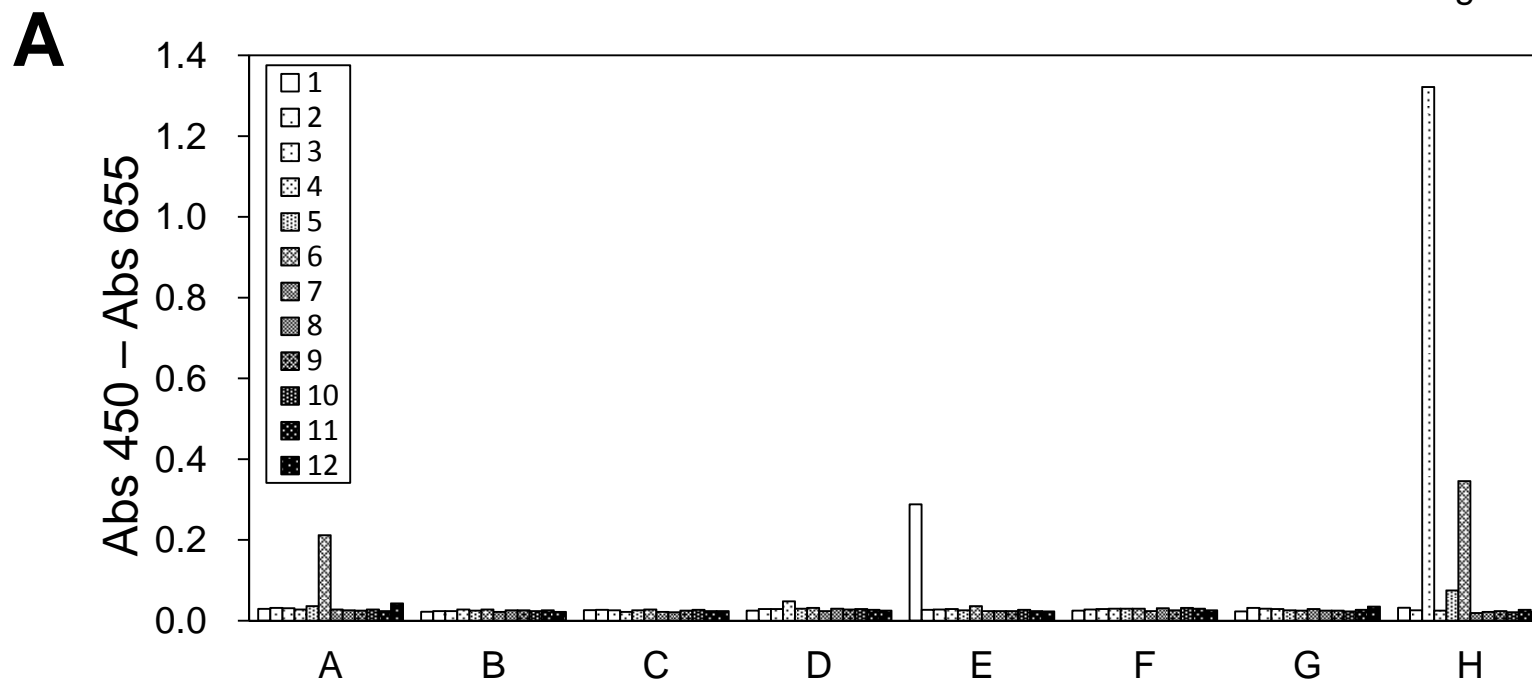
444

1 **TABLE 1.** Kinetic parameters of the monoclonal antibodies

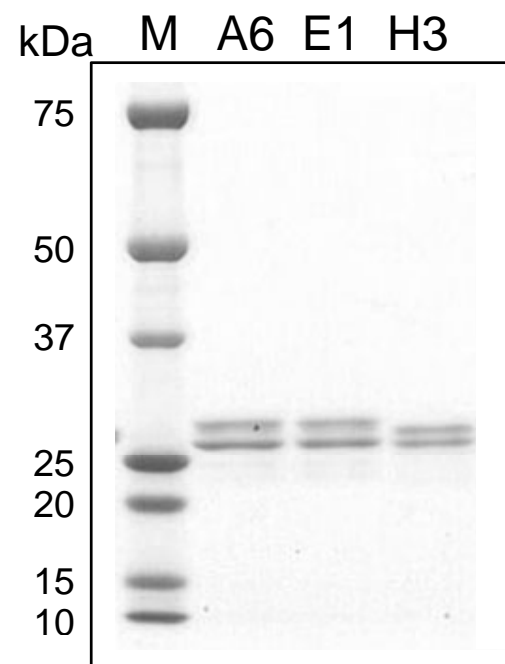
Clone name	K_a ($10^4/\text{Ms}$)	K_d ($10^{-3}/\text{s}$)	K_D (nM)
A6	5.80	24.0	420
E1	0.57	29.0	5100
H3	1.20	0.87	70

2

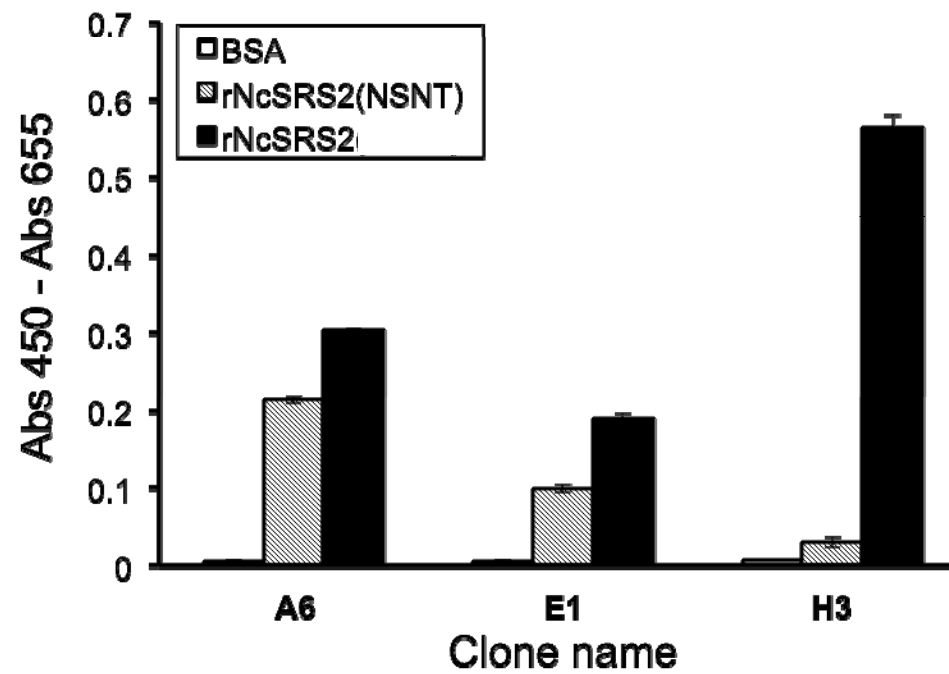


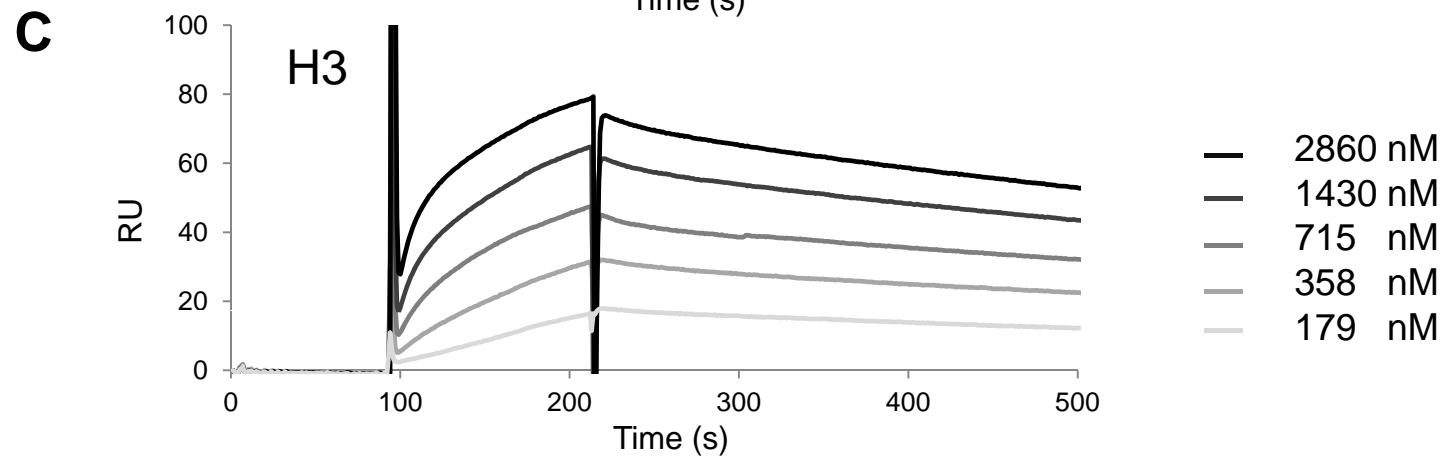
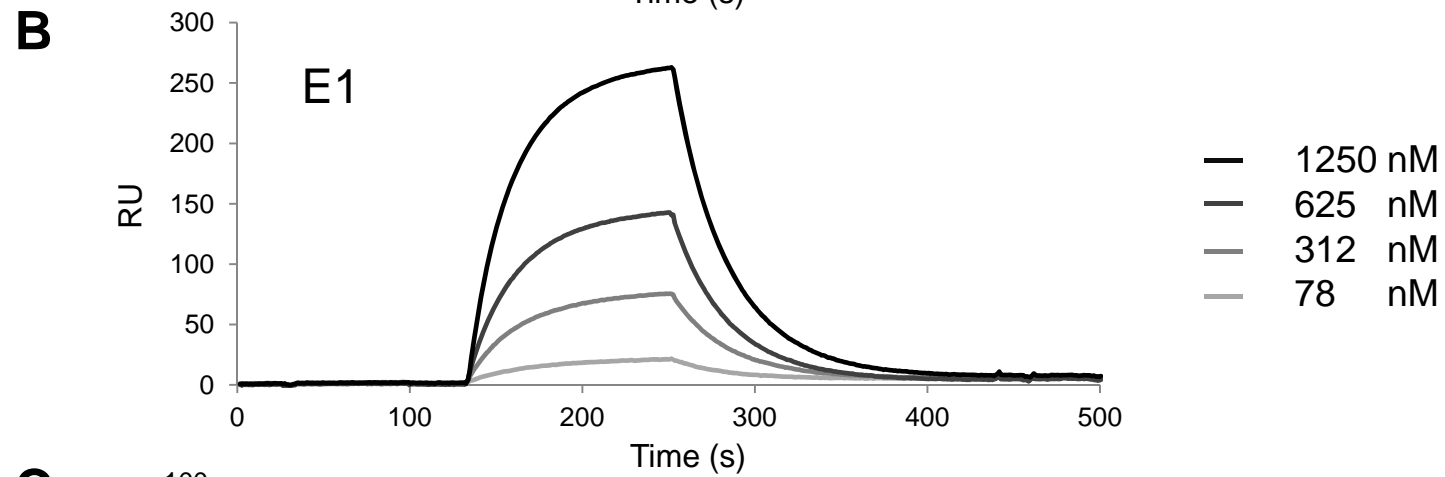
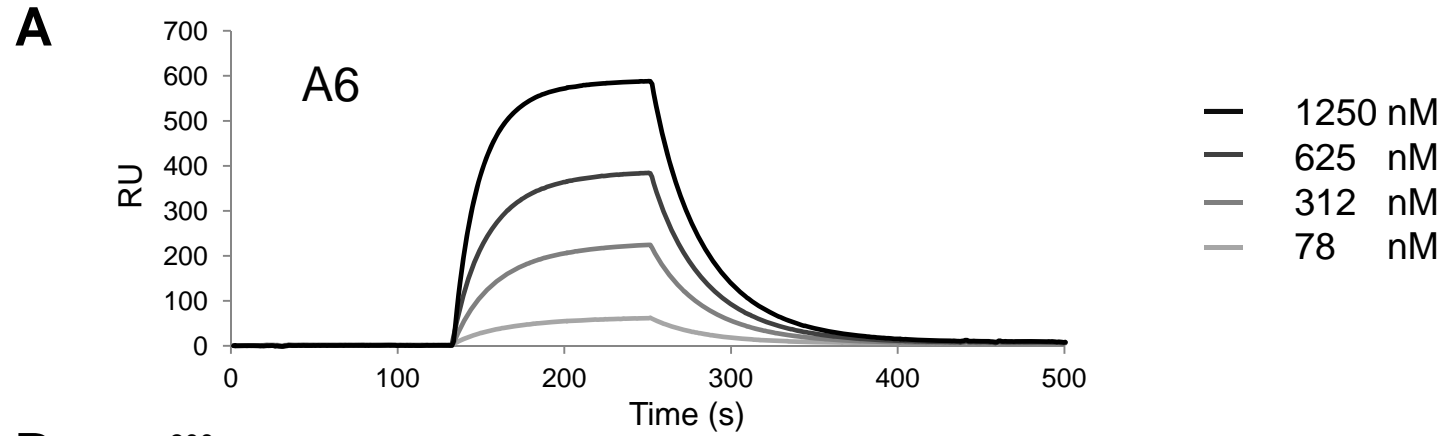


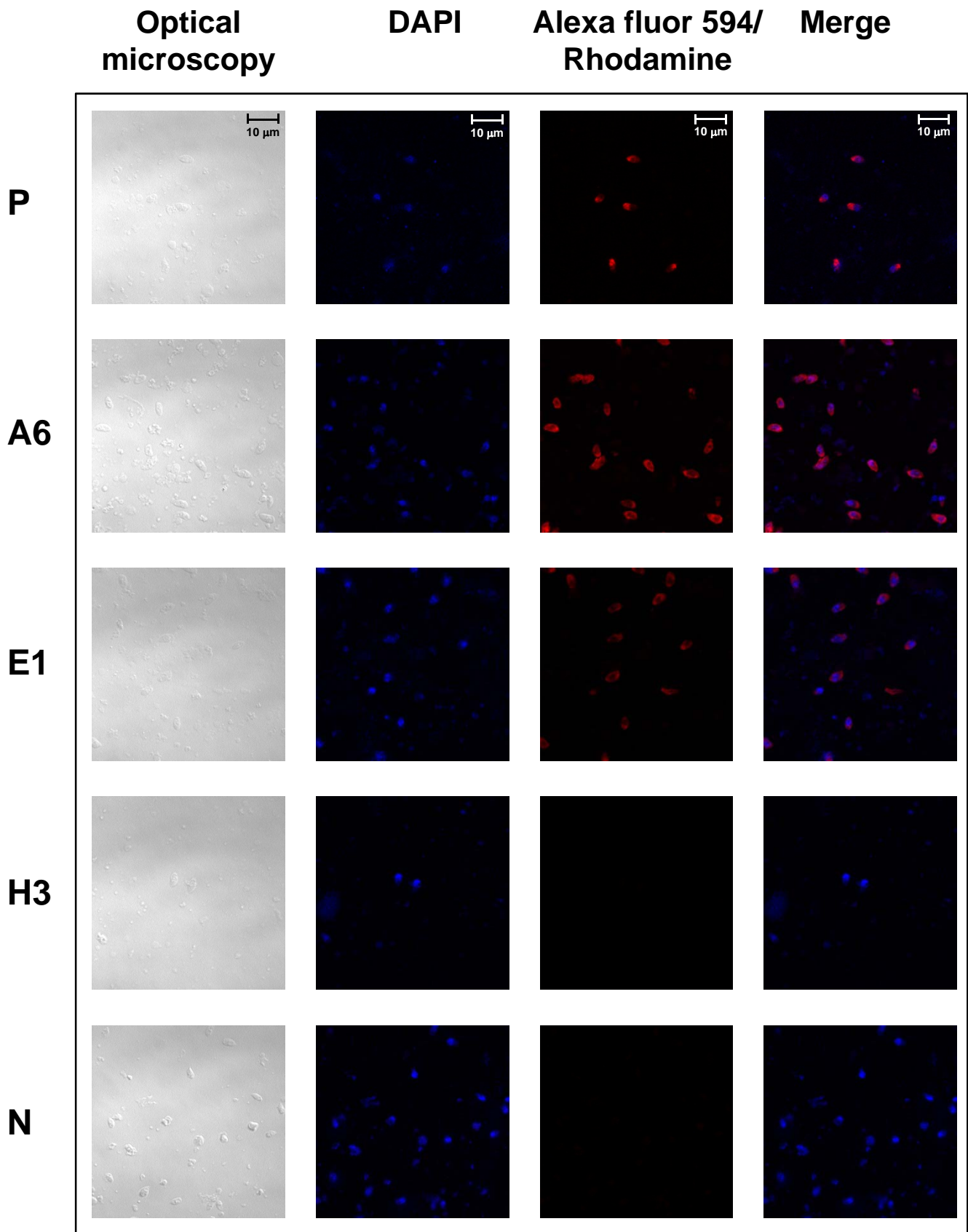
A



B







		CDR-L1		CDR-L2				
A6VL	1	ENVLTQSPA	IMSASPGEKVTMTCT	SASSSVS-----YMH	WYQQKSGTSPKRWIY	DTSKLAS	GVPARFSG 70	
E1VL	1	ENVLTQSPA	IMSASPGEKVTMTCT	SASSSVS-----YMH	WYQQKSGTSPKRWIY	DTSKLAS	GVPARFSG 70	
H3VL	1	DIVMTQSQK	FMSTSVGGRVSITCT	KASQSVRT-----AVA	WFQQKSGQSPEALIY	LASNRHT	GVPDRFTG 70	
		CDR-L3						
	71	SGSGTSYSLT	ISSMEAEDAATYYC	QQWNN--IPPT	FGAGTKLELKRA		117	
	71	SGSGTSYSLT	ISSMEAEDAATYYC	QQWNN--IPPT	FGAGTKLELKRA		117	
	71	SGSGTDFTLT	ISNVQSEDLADYFC	LQHWN--YPYT	FGGGTKLEIKRA		117	
		CDR-H1		CDR-H2				
A6VH	118	QVQLKESGGGLV	KPGGSLKLSCAAS	GFTFSS--YAMS	WVRQTPEKRLEWVA	SISSGG---STY	YPDSV 185	
E1VH	118	E V K L V E	ESGGGLV	KPGGSLKLSCAAS	EFTFSS--YAMS	WVRQTPEKRLEWVA	SISSGG---STY	YPDSV 185
H3VH	118	QVHLQQSGAELAK	PGASVKMSCKAS	GYTFTS--YWMH	WVKQRPGQGLEWIG	YINPSTG--YTE	YNQKF 185	
		CDR-H3						
	186	KGRFTISRDNARN	IYLQMSLRS	EDTAMYICAR	RGGLG-----FDY	WGQGTTLTVS	249	
	186	KGRFTISRDNARN	IYLQMSLRS	EDTAMYICAR	RGGLG-----FDY	WGQGTTLTVS	249	
	186	KDKATLTADKSS	STAYMQLSSLT	SEDSAVYYCAR	EY-----FDY	WGQGTTLTVS	249	