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Evaluating the 3C-like protease activity of SARS-Coronavirus: Recommendations for standardized assays for drug discovery

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Abstract

Although the initial outbreaks of the deadly coronavirus that causes severe acute respiratory syndrome (SARS-CoV) were controlled by public health measures, the development of vaccines and antiviral agents for SARS-CoV is essential for improving control and treatment of future outbreaks. One potential target for SARS-CoV antiviral drug development is the 3C-like protease (3CLpro). This enzyme is an attractive target since it is essential for viral replication, and since there are now a number of high resolution X-ray structures of SARS-CoV 3CLpro available making structure-based drug-design possible. As a result, SARS-CoV 3CLpro has become the focus of numerous drug discovery efforts worldwide, but as a consequence, a variety of different 3CLpro expression constructs and kinetic assays have been independently developed making evaluation and comparison between potential inhibitors problematic. Here, we review the literature focusing on different SARS-CoV 3CLpro expression constructs and assays used to measure enzymatic activity. Moreover, we provide experimental evidence showing that the activity of 3CLpro enzymatic is significantly reduced when non-native sequences or affinity-tags are added to the N- or C-termini of the enzyme, or when the enzyme used in assays is at concentrations below the equilibrium dissociation constant of the 3CLpro dimer. We demonstrate for the first time the utility of a highly sensitive and novel Alexa488-QSY7 FRET-based peptide substrate designed for routine analysis and high-throughput screening, and show that kinetic constants determined from FRET-based assays that are uncorrected for inner-filter effects can lead to artifacts. Finally, we evaluated the effects of common assay components including DTT, NaCl, EDTA and DMSO on enzymatic activity, and we recommend standardized assay conditions and constructs for routine SARS-CoV 3CLpro assays to facilitate direct comparisons between SARS-CoV 3CLpro inhibitors under development worldwide.

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1. Introduction

Severe acute respiratory syndrome (SARS) is a form of atypical pneumonia first identified in late 2002. The disease was found to be caused by a novel human coronavirus named SARS coronavirus (SARS-CoV) in early 2003 (Drosten et al., 2003; Ksiazek et al., 2003; Peiris et al., 2003). Coronaviruses are enveloped, positive-stranded RNA viruses containing the largest viral RNA genomes known to date (27–31 kb). These RNA coronaviruses release their genetic information into the cytoplasm of the cell. The S'-two thirds of the SARS-CoV genomic RNA contains two open reading frames (ORFs), termed ORF1a and ORF1b. These ORFs are translated via a ribosomal frameshift mechanism (Thiel et al., 2003) to generate the viral replicase polyproteins pp1a and pp1ab that mediate the functions required for viral replication and transcription (Marra et al., 2003; Rota et al., 2003). These polyproteins are extensively processed by viral proteases to generate 16 nonstructural proteins (nsps) (Fig. 1) that assemble with host intracellular membranes to form double membrane vesicles (DMVs), the site of viral RNA synthesis (Goldsmith et al., 2004; Gosert et al., 2002; Snijder et al., 2006).

Functional polypeptides are released from the pp1a and pp1ab polyproteins via extensive proteolytic processing by the papain-like proteinase (PLpro) and the 3C-like protease...
Fig. 1. Schematic diagram of the translated SARS-CoV genomic RNA in which the open reading frames encoding the viral replicase polyproteins, pp1a and pp1ab, are depicted. The 3 sites recognized and processed by PLpro (red) and the 11 sites recognized and processed by 3CLpro (blue) are indicated. Complete processing of the 14 sites produces 16 non-structural proteins (nsp’s). Transmembrane regions are indicated as “TM”, and the polymerase (Pol) and helicase (Hel) coding regions are indicated.

(3CLpro), which are located in the non-structural protein (nsp) regions nsp3 and nsp5, respectively (Fig. 1). Both proteases are processed autocatalytically from the replicase polyprotein sequence, and PLpro is responsible for cleaving 3 sites whereas 3CLpro is responsible for cleaving 11 sites within the viral genome (Harcourt et al., 2004; Thiel et al., 2003). The enzymatic activity of both PLpro and 3CLpro are essential for the viral life cycle, and therefore both enzymes are attractive targets for the development of antiviral drugs directed against SARS-CoV and other coronavirus infections.

To date, the overwhelming majority of research groups worldwide are focusing their drug-discovery efforts on the 3CLpro enzyme. This is most likely due to the facts that; (1) the closely related 3C-like proteinase from the human rhinovirus has been successfully targeted by antiviral agents for treatment of the common cold (Matthews et al., 1999); (2) there are a number of X-ray structures available for the 3C-like proteases from SARS-CoV, rhinoviruses, HCoV-229E, and transmissible gastroenteritis virus (TGEV) making structure-based drug design feasible (Anand et al., 2002, 2003; Ghosh et al., 2005; Hilgenfeld et al., 2006; Matthews et al., 1999); and (3) the SARS-CoV 3CLpro enzyme can be purified in large quantities, is highly active and easily assayed, and is readily crystallized making in vitro studies very straightforward. In contrast, the SARS-CoV PLpro enzyme is membrane associated making it significantly more challenging to study in vitro. Recently, a soluble form of

Fig. 2. SARS-CoV 3CLpro cleavage sites and the canonical recognition sequence. The 11 recognition sequences of SARS-CoV 3CLpro from the P6 to P5’ positions are shown with their respective locations between the respective nsp’s. The relative kcat/Km values for the series of 11-mer peptide substrates are also given (Fan et al., 2005). A canonical recognition sequence is proposed based on the relative kcat/Km values for SARS-CoV 3CLpro, and the recognition sites of a series of other coronavirus 3C proteases (Kiemer et al., 2004; Thiel et al., 2003).
SARS-CoV PLpro has been overexpressed, purified and characterized (Barretto et al., 2005, 2006), and its’ X-ray structure has been determined by our group (Ratia et al., 2006).

So far, only a handful of compounds have been synthesized that exhibit submicromolar IC50 values against SARS-CoV 3CLpro activity in vitro, and the majority of these compounds either do not have antiviral activity, or have not been evaluated in vivo. The discovery and development of antiviral compounds is, in general, coupled to an in vitro enzymatic assay that can be performed in a laboratory setting. Frequently, as is the case for 3CLpro, these assays are based on hydrolysis of a small synthetic peptide which can be monitored by either HPLC analysis, or by monitoring the release of a chromophore or fluorescent reporter group upon cleavage. For SARS-CoV 3CLpro, there have been a number of studies that have each utilized different peptide substrates, as well as different reporter groups including chromophores, fluorescence groups, and fluorescence resonance energy transfer (FRET) pairs (Table 1). All three reporters have distinct advantages over HPLC detection since they are typically more sensitive and offer a methodology for continuous monitoring of proteolytic activity over time.

A variety of substrates, based on the 3CLpro recognition sequence (Fig. 2), have been used in the FRET-based assays (Table 1) (Bacha et al., 2004; Graziano et al., 2006; Kao et al., 2004; Kuo et al., 2004). Moreover, these studies have each utilized a variety of assay conditions and different SARS-CoV 3CLpro constructs that do not necessarily reflect the authentic sequences in vivo. Evidence is mounting that the kinetic properties of 3CLpro are significantly influenced by the construct and assay conditions utilized (Table 2). Tied to this knowledge is the observation that SARS-CoV 3CLpro is only functional as a dimer in solution (Chen et al., 2005a; Fan et al., 2004; Shi et al., 2004). Therefore, it is important to understand these differences and to develop a set of standardized assay conditions containing components appropriate for comparing differences in kinetic properties and parameters, e.g. IC50 and Ki values.

| Donor, fluorophore, or colorimetric label | Quencher | Wavelength (Ex/Em) | P7 P6 P5 P4 P3 P2 P1 P1' P2' P3' P4' P5' P6' P7' | References |
|------------------------------------------|----------|--------------------|-------------------------------------------------|------------|
| EDANS                                    | Dabcyl   | 330-360 nm/490-538 nm | E* S A T L Q S G L A K* S K* T S A V L Q S G F R K K* E* | This work |
| Alexa Fluor 488                          | QSY 7    | 490 nm/535 nm       | C* E S A T L Q S G L A K* | This work |
| Alexa Fluor 594                          | QSY 21   | 595 nm/620 nm       | C* E S A T L Q S G L A K* | This work |
| Abz                                      | nitrotyrosine | 320-340 nm/400-440 nm | S* V T I L Q S G V* R | This work; (Blanchard et al., 2004; Kaeppler et al., 2005; Lee et al., 2005) |
| Abz                                      | Dnp      | 320-362 nm/420-425 nm | S* A V L Q S G F R K* T* S A V L Q S G F R K* S* G V T F Q G K F K K* | (Liu et al., 2005a) |
| mCA                                      | Dnp      | 320 nm/405 nm       | A* V L Q S G F K* K | (Yang et al., 2005) |
| Rhodamine (fluorophore)                  |          | 492 nm/523 nm       | A R L Q* | (Graziano et al., 2006) |
| pNA (colorimetric label)                 |          | 390 nm              | T S A V L Q* | (Chen, Wei et al., 2006; Huang et al., 2004; Liu et al., 2005b; Wei et al., 2006) |

*Attachment site of donor, fluorophore, or colorimetric label.

*Attachment site of quencher label.
between different compounds aimed at serving as therapeutic leads. In the occurrence of drug design, it becomes particularly vital to standardize an approach that can be used by the scientific community for reliable comparison of different or even the same compounds between research groups.

In this study, we compared the enzymatic activity between two different constructs of SARS-CoV 3CLpro using FRET-based assays, and we evaluated the influence of reducing agents, salts, EDTA and DMSO, which is often used as a solvent to dissolve inhibitors, on SARS-CoV 3CLpro enzyme activity. In addition, we compared the activity and sensitivity of four different FRET-based peptide-substrates. The resulting data allow for the recommendation of a standard set of assay conditions and SARS-CoV 3CLpro constructs that can be used and universally adopted.

2. Materials and methods

2.1. Plasmid construction

The gene encoding 3CLpro (SARS-CoV polyprotein residues 3241–3544) was amplified using SuperScript One-Step RT-PCR for Long Templates (Invitrogen) according to the manufacturer’s instructions. RNA isolated from SARS-CoV-(Urbani strain)-infected VeroE6 cells served as template for the RT-PCR reaction. The forward and reverse primers were as follows: 5’-GGTCTAGCTAGCGGTTTTAGGAAAATGGCATTC-3’ and 5’-CCTAGCTCAGCTTAGGTAACACCAGAGCATTGTC-3’, respectively. PCR product was digested with the restriction enzymes NheI and Bpu1102I and ligated into similarly digested pET11a vector (Novagen). The resulting plasmid DNA was designated pET11a-SCoV-3CLpro. To generate the 6xHis-tagged construct, the 3CLpro sequence was amplified using forward primer 5’-GGTCTACTCGAGAGTGGTTTTAGGAAAATGGCA-3’ and the above reverse primer. The resulting PCR product was digested with XhoI and Bpu1102I ligated into pET15b, and designated pet15b-SCoV-3CLpro. Constructs were verified by DNA sequencing.

2.2. Expression and purification of SARS-CoV 3CLpro

3 L of E. coli BL21(DE3) cells, transformed with pET11a-3CLpro plasmid, were grown for 24 h at 25 °C without induction. Cells were pelleted by centrifugation and resuspended in 100 mL of Buffer A [20 mM Tris pH 7.5, 2.5 mM dithiothreitol (DTT)] containing 500 μg of lysozyme. The cells were incubated for 10 min on ice and then lysed via sonication using a 600-watt Model VCX ultrasonicator. After pelleting the cell debris by centrifugation (40,900 × g for 30 min), the clarified cell lysate was loaded onto a 120 mL DEAE Sepharose Fast Flow column (Amersham Biosciences, Piscataway, NJ) equilibrated with Buffer A. Since 3CLpro does not bind to DEAE resin at pH...
7.5, column effluent containing 3CLpro was collected and subjected to a 60% ammonium sulfate fractionation. The suspension was centrifuged (40,900 × g for 30 min) and the resulting pellet was resuspended in 1 M ammonium sulfate, 20 mM Tris pH 7.5, 2.5 mM DTT. The dissolved pellet was loaded onto a 50 mL Phenyl Sepharose 6 Fast Flow HS column (Amersham Biosciences) equilibrated with Buffer B (1.5 mM ammonium sulfate, 20 mM Tris pH 7.5, 2.5 mM DTT). Protein was eluted with a 10× column volume gradient to 100% Buffer A. Fractions containing 3CLpro were pooled, concentrated, and diluted 30-fold with Buffer A before being loaded onto a Mono Q 10/10 column (Amersham Biosciences) equilibrated in the same buffer. A 10× column-volume gradient from 0–100% Buffer C (20 mM Tris pH 7.5, 0.5 M NaCl, 0.5 M imidazole, 2.5 mM DTT) was used to elute the protein. Pure 3CLpro fractions were exchanged into Buffer A containing 10% glycerol and concentrated to approximately 40 mg/mL for use in kinetic assays and crystallization experiments.

SARS-CoV 3CLpro with an N-terminal (His)6-tag was purified from BL21(DE3) cells expressing the pET15b construct. 1 L of cells were grown for 24 h at 25 °C without induction. Pelleted cells were resuspended in 40 mL of Buffer D (20 mM Tris pH 7.5, 0.5 M NaCl, 10 mM imidazole, 2.5 mM DTT) and were lysed as above. The clarified lysate was applied to a 5 mL HisTrap column (Amersham Biosciences) which was equilibrated with Buffer D. Protein was eluted from the column with a 20 column-volume gradient to 100% Buffer A. Fractions containing pure protein were pooled, concentrated, and exchanged into Buffer A containing 10% glycerol and concentrated to approximately 40 mg/mL for use in kinetic assays and crystallization experiments.

SARS-CoV 3CLpro with an N-terminal (His)6-tag was purified from BL21(DE3) cells expressing the pET15b construct. 1 L of cells were grown for 24 h at 25 °C without induction. Pelleted cells were resuspended in 40 mL of Buffer D (20 mM Tris pH 7.5, 0.5 M NaCl, 10 mM imidazole, 2.5 mM DTT) and were lysed as above. The clarified lysate was applied to a 5 mL HisTrap column (Amersham Biosciences) which was equilibrated with Buffer D. Protein was eluted from the column with a 20 column-volume gradient to 100% Buffer A. Fractions containing pure protein were pooled, concentrated, and exchanged into Buffer A containing 10% glycerol and concentrated to approximately 40 mg/mL for use in kinetic assays and crystallization experiments.

2.4. Determination of fluorescence extinction coefficients

The amount of product produced over time was calculated from a fluorescence extinction coefficient (FEC) that was determined for each FRET peptide-substrate under a given assay and instrument settings. A set of 100 µL reactions were performed in triplicate using 50 mM HEPES buffer, pH 7.5, and fluorogenic substrates concentrations of 0.0, 0.1, 0.5, 1.0, 2.0, and 5.0 µM. Excess 3CLpro enzyme was added to rapidly cleave 100% of the substrate. The FEC values were then determined from a standard curve whereby the concentration of substrate was plotted against the total arbitrary fluorescence produced from a standard curve whereby the concentration of substrate minus the AFU values produced from a control with no enzyme added. All data were also corrected for any inner-filter effect caused by the samples absorbing too much light which was observed for all substrates tested (Liu et al., 1999). The resulting data were then fit to a line and the resulting slope is designated the fluorescence extinction coefficient (AFUMAX/µM) of the respective fluorogenic substrate.

2.5. SARS-CoV 3CLpro reaction rates and turnover numbers

All reactions were performed in duplicate or triplicate to determine the initial reaction rates. The reaction volumes were 100 µL in 50 mM HEPES buffer, pH 7.5, using a fluorogenic substrate concentration of 1 µM. Baseline hydrolysis rates were also measured using reactions without enzyme. Initial rates were determined by monitoring the change in AFUs per min,
which were then converted to the amount of product produced per minute in μM/min using the fluorescence extinction coefficient. Turnover numbers (min⁻¹) were then calculated from the molecular weights of the 3CLpro constructs.

2.6. Influence of NaCl, DTT, EDTA and DMSO on SARS-CoV 3CLpro activity

The influence of assay components on SARS-CoV 3CLpro activity was measured using the AlexaFluor® 488-QSY®-7 FRET peptide substrate. All reactions were performed in triplicate to determine the effect of sodium chloride [NaCl], dithiothreitol [DTT], ethylenediaminetetraacetic acid [EDTA], and dimethyl sulfoxide [DMSO] on enzyme activity. One hundred μL reactions containing HEPES buffer [50 mM, pH 7.5] and 1 μM fluorogenic substrate were supplemented with either NaCl (0–1 M), DTT (0–10 mM), DMSO (0–10%) and EDTA (0–2 mM). Tagged or untagged 3CLpro enzyme was added to a final concentration of 100 nM to initiate the reaction.

2.7. Influence of dimerization on SARS-CoV 3CLpro activity

The activity of the native and (His)₆-tagged SARS-CoV 3CLpro constructs as a function of enzyme concentration were measured in triplicate over a concentration range of 50–1000 nM using the AlexaFluor® 488-QSY®-7 FRET peptide substrate. The reaction rate data were then plotted as a function of 3CLpro enzyme concentration and fit to Eq. (1) that describes the rate of an active dimeric enzyme that is in equilibrium with an inactive monomer.

\[
V_{\text{max}} = K_{\text{cat}}[D] = K_{\text{cat}} = \frac{K_d + 4C_T - \sqrt{K_d^2 + 8K_dC_T}}{8} \quad (1)
\]

Here, \(K_d\) is the apparent dissociation constant of the dimer and is given by \(K_d = [M][M]/[D]\) where [M] and [D] are the concentrations of the monomer and dimer. \(C_T\) is the total concentration calculated as the monomer, and is related to the dimer concentration by \(C_T = [M] + 2[D]\). The data were fit using non-linear regression to different \(C_T\) values using Eq. (1) and the Program TableCurve2D (SPSS Scientific).

3. Results and discussion

3.1. Purification of native and (His)₆-tagged SARS-CoV 3CLpro and evaluation of new FRET-peptide substrates with AlexaFluor dyes and QSY quenchers

Native and (His)₆-tagged versions of SARS-CoV 3CLpro were expressed and purified from E. coli BL21(DE3) cells. The native version of the enzyme is the exact primary sequence that would be released from the SARS-CoV polyprotein with the addition of a N-terminal methionine residue. Native 3CLpro was purified to homogeneity in four steps including weak anion-exchange filtration (DEAE), ammonium sulfate fractionation, hydrophobic interaction (phenyl sepharose) and anion-exchange (Mono-Q) chromatography. The yield of enzyme from 31 is approximately 100 mg, and the final enzyme is over 95% pure as judged by SDS-PAGE analysis (Fig. 3). Purification of the (His)₆-tagged enzyme was achieved in a single step with a yield of approximately 50 mg from 11, and the enzyme is over 95% pure by SDS-PAGE analysis (Fig. 3).

In order to measure the activity of the SARS-CoV 3CLpro enzyme, we evaluated a series of FRET-based peptides for their ability to serve as substrates. The primary sequence of these substrates is based on the canonical peptide sequence shown in Fig. 2. Two of the FRET reporter groups, Dabcyl-EDANS and Abz-Tyr(NO₂), were chosen because they are readily synthesized commercially, and they are commonly used for the measurement of protease activity. For SARS-CoV 3CLpro, Dabcyl-EDANS has been the most popular reporter system (Chen et al., 2005a,b,c; Wu et al., 2004, 2006) (Table 1). Other groups have utilized Abz or MCA fluorophores with the corresponding nitrotyrosine or Dnp quenchers (Kaeppler et al., 2005; Kuang et al., 2005; Lee et al., 2005; Yang et al., 2005) in addition to colorimetric labels (Liu et al., 2005b) (Table 1). Although these FRET-pairs have an established history, there are a number of newer fluorophores on the market that have higher quantum yields, pH independent fluorescence, and absorbance spectra in the visible region (e.g. >450 nm) making them potentially useful for high-throughput screening and ultra-sensitive fluorescence measurements. One series of fluorophores are the AlexaFluor dyes (Molecular Probes & Invitrogen).

The AlexaFluor dyes are extremely fluorescent, pH insensitive, water soluble and highly stable, and are therefore commonly used in flow cytometry, immunocytochemistry, and immunohistochemistry experiments. Despite their many potential advantages, they have not been utilized in FRET-peptide substrates for protease assays. Therefore, we evaluated two of the AlexaFluor dyes, AlexaFluor® 488 and AlexaFluor® 594, in combination with QSY quenchers QSY®-7 and QSY®-21.
resulted from a fit of the data to a line using linear regression. The slope of the line that was fit to the data in Fig. 4. The slope, or FEC value, values and their errors are listed in Table 3.

The concentration of FRET-peptide was varied and plotted against the total arbitrary fluorescence values (AFUs) released upon complete cleavage of 1 μM of substrate by the native, i.e. non-tagged, enzyme. The results are summarized in Table 3 and indicate that the Alexa488-QSY7, Dabcyl-EDANS, and Abz-Tyr(NO2) substrates all efficiently catalyzed the cleavage reaction at 0.028 to 0.068 min⁻¹, whereas the Alexa594-QSY21 peptide substrate is cleaved approximately 10 times more slowly. Since the size of the Alexa Fluo 594 dye is significantly larger than the other fluorophores (Table 1), some steric blockage may result and thereby decrease the catalytic rate with this substrate.

Each combination of FRET groups has different advantages and disadvantages depending on the required application. For instance, significantly increased sensitivity can be obtained by using the Alexa488 dye which has an excitation wavelength in the visible range that is beyond most small molecules. These properties make this dye particularly attractive for conducting high-throughput screening studies. In addition, the sensitivity allows the measurement of routine IC₅₀ values at substrate concentrations significantly below the Kₘ values which saves on substrate and makes the IC₅₀ values closer to the correct Kᵢ values.

A disadvantage of highly fluorescence FRET-substrates such as Alexa-QSY and Dabcyl-Edans, is that they are susceptible to a phenomenon known as the “inner-filter effect”. This effect results when the fluoresced light is absorbed by quenching groups on neighboring substrates or cleaved product molecules so that only a fraction of the fluoresced light impinges upon the detector system of the fluorimeter (Liu et al., 1999). The result of the inner-filter effect is that if uncorrected, incorrect or artificial kinetic parameters, i.e. Kₘ and Kᵢ, are determined. Liu and co-workers have demonstrated that this is a common problem with Dabcyl-EDANS FRET-substrates and depends on the wavelength range, pathlength, and concentration of quenching components (Liu et al., 1999). A survey of the literature on the influence of the inner-filter effect on SARS-CoV 3CLpro assays utilizing Dabcyl-EDANS and Abz-Tyr(NO2) substrates produced a small number of studies that observed and corrected for the inner-filter effect (Blanchard et al., 2004; Kaepppler et al., 2005; Lee et al., 2005), and a larger number of studies that did not account for this effect or report it (Bacha et al., 2004; Chou et al., 2004; Kao et al., 2004; Kuo et al., 2004; Liu et al., 2005a). When the inner-filter effect is corrected, the Kₘ values of FRET-peptide substrates range from 190 to 2 mM depending on the peptide sequence, with the majority of Kₘ values greater than 500 μM (Blanchard et al., 2004; Fan et al., 2004; Kaepppler et al., 2005; Lee et al., 2005). These values are commensurate with the Kₘ values derived from HPLC assays. In contrast, the Kₘ values derived uncorrected data are typically in the 10–50 μM range (Bacha et al., 2004; Chou et al., 2004; Kao et al., 2004; Kuo et al., 2004; Liu et al., 2005a). What is most revealing in a number of these studies is that the uncorrected FRET-peptide substrate Kₘ data are different than the HPLC derived kinetic data. Kao and co-workers show a plot of uncorrected rate data versus Dabcyl-EDANS, FRET-substrate concentration with apparent substrate inhibition noticeable at 40 μM (Kao et al., 2004) which is exactly what is observed for uncorrected data (Liu et al., 1999). We observed and corrected for the inner-filter effect with all four substrates tested.

| FRET-peptide substrate | FEC (AFU/μM) | Activity (min⁻¹) |
|------------------------|--------------|------------------|
| Alexa488-QSY7          | 37,840 ± 15  | 0.038            |
| Alexa594-QSY21         | 3,350 ± 100  | 0.003            |
| Dabcyl-Edans           | 4,530 ± 150  | 0.068            |
| Abz-Tyr(NO2)           | 830 ± 50     | 0.028            |

* The specific sequence for each FRET-peptide substrate is described in Section 2.
* The fluorescence extinction coefficient values (FECs) were determined from the slope of the line that resulted from a fit of the data to a line using linear regression.
* The activity, i.e. turnover number was determined at a substrate concentration of 1 μM and a non-tagged enzyme concentration of 100 nM.

Fig. 4. Fluorescence extinction coefficient (FEC) determination of four FRET-peptide substrates. The concentration of FRET-peptide was varied and plotted against the total arbitrary fluorescence values (AFUs) released upon complete cleavage of the substrate by an excess of SARS-CoV 3CLpro. Data were fit to a line to a line and the slope is the FEC constant in units of AFUs/μM. The FEC values and their errors are listed in Table 3.
A wide range of (His)6-tagged SARS-CoV 3CLpro is approximately 30-times apparent from the plots in Fig. 5 is that the activity of the enzyme at fixed substrate concentrations of 1 mM results in best-fit parameters for Alexa488-QSY7 were 3.2. Influence of (His)6-affinity tags on 3CLpro activity and crystallization

A survey of the available literature on the SARS-CoV 3CLpro constructs used for expression and purification indicate a wide range of constructs as well as methods used to produce the final purified enzymes (Chen et al., 2005a, 2006; Chou et al., 2004; Fan et al., 2004; Hsu et al., 2005; Kuo et al., 2004; Shi and Song, 2006; Shi et al., 2004; Sun et al., 2003; Wei et al., 2006). A number of the constructs used and reported in the literature are listed in Table 2 along with their relative activities. We noticed a wide range of $K_i$ values and turnover numbers reported for different constructs, and had a difficult time ascertaining whether or not the differences in values for these kinetic constants were due to the differences in the assay conditions (addressed below), or due to differences in the SARS-CoV 3CLpro constructs used in the studies. Therefore, we tested directly the influence of a (His)6-tag on the catalytic activity of SARS-CoV 3CLpro. The enzymatic activity of native and n-(His)6-tagged SARS-CoV 3CLpro constructs as a function of enzyme concentration for two different FRET-peptide substrates, Alexa488-QSY7 and Dabcyl-EDANS, are shown in Fig. 5. The assays were performed at fixed substrate concentrations of 1 μM. What is immediately apparent from the plots in Fig. 5 is that the activity of the n-(His)6-tagged SARS-CoV 3CLpro is approximately 30-times lower than the native enzyme for both substrates tested. Moreover, at a concentration of 1 μM enzyme, the n-(His)6-tagged construct still lags far behind in activity compared to the native enzyme.

The response of the native enzyme activity to increasing concentrations of enzyme over the range of 50–1000 nM is non-linear especially at lower enzyme concentrations. This curvature indicates that dimer-to-monomer dissociation may be occurring. A number of studies have demonstrated that SARS-CoV 3CLpro functions as a dimer (Chen et al., 2005a; Fan et al., 2004; Shi et al., 2004). We have compiled a list of different dimer-to-monomer dissociation constants in conjunction with the various constructs and methods used to measure the oligomeric state in Table 2. We fit our data in Fig. 5 to Eq. (1) which describes dimer-to-monomer dissociation where only the dimer is active (Chen et al., 2006; Kuo et al., 2004). From a fit of our data to Eq. (1), we obtain dissociation constants between 0.25 and 1 μM considering the error in the fits. Our values are commensurate with the enzymatic-derived values of Chen and coworkers for an uncleaved, c-(His)6-tagged construct (Chen et al., 2006; Kuo et al., 2004), lower than the values of Fan and coworkers for an uncleaved, c-(His)6-tagged construct (Fan et al., 2004), and higher than the values obtained by Kuo and coworkers for a cleaved n-(His)6-tagged construct (Kuo et al., 2004). In general, our values are consistently and significantly lower than the dissociation constants for the tagged-constructs that were determined by any of the biophysical methods (Table 2).

As can be seen from Table 2, the trend toward using 3CLpro-constructs with modified N- and C-terminal ends shifts the equilibrium towards the monomer resulting in higher dissociation constants. Although investigators have claimed the use of (His)6-tags have no effect on enzymatic or quaternary structure (Bacha et al., 2004; Chou et al., 2004), it is clear that in our experiments, N-terminal tags have a significant effect on kinetic parameters (Fig. 5). The influence of N- and C-terminal activity can be explained partially by the nature of dimerization of the enzyme structure. The interactions of the N- and C-terminal extensions of monomers of SARS-CoV 3CLpro near the active site are shown in Fig. 6. This figure was generated from the X-ray crystal structure of the Cys145Ala mutant of SARS-CoV 3CLpro (PDB 1Z1J: (Hsu et al., 2005)) that has small N- and C-terminal extensions. In this structure, the C terminus of a protomer from another asymmetric unit is intercalated into the active site of the 3CLpro dimer. The C-terminal peptide containing a glutamine residue occupies the P1 pocket of the enzyme. The N-terminal peptide from the other monomer of the dimer is modeled in yellow and contains the serine residue in the P1’ pocket of the enzyme. This is representative of the autoprocessing cleavage reaction that occurs in vivo. Note that the ends...
of the molecules extend directly into the active site. Thus, the introduction of tags on the N- or C-termini, will likely influence the quaternary structure and hence enzymatic properties of the enzyme. In strong support of our observations, Xue and coworkers reported recently that the addition of two residues (Gly-Ser) or five residues (Gly-Pro-Leu-Gly-Ser-) to the N-terminus of SARS-CoV 3CLpro reduced its catalytic efficiency ($K_{cat}/K_m$) by 20-fold and 150-fold, respectively (Xue et al., 2007).

We also tested the influence of the (His)$_6$-tag on the crystallization of SARS-CoV 3CLpro. We were unable to crystallize the apoenzyme or covalently modified enzyme from over 1000 crystallization conditions attempted. In contrast, we can readily crystallize the covalently labeled native enzyme within 4 h (Fig. 3), and the apoenzyme within a couple of days. The negative influence of the N-terminal extension on crystallization also supports the kinetic and other biophysical data suggesting a profound influence of N- and C-terminal extensions on the properties of SARS-CoV 3CLpro. Therefore, our experimental observations, coupled with the vast amount of literature now available on the quaternary structure of SARS-CoV 3CLpro, leads us to conclude that native constructs are much preferred over uncleaved, tagged constructs for analysis of 3CLpro enzymatic activity and structure.

### 3.3. Influence of assay components on 3CLpro activity

In constructing the summaries of 3CLpro substrates and properties presented in Tables 1 and 2, we also noted a significant variation in the buffer components and concentrations used in SARS-CoV 3CLpro assays. Variations included buffer type and pH, type and concentration of salt, reducing agents such as DTT and BME, metal chelating agents such as EDTA, detergents, DMSO, and BSA. Since the addition of one or a combination of these components can have a significant influence on activity, we determined their influence on the activity of native SARS-CoV 3CLpro using the Alexa488-QSY7 peptide-substrate. We chose a pH value of 7.5 since the overwhelming amount of data available on the pH rate profiles for the enzyme suggest a pH optima at this value (Chen et al., 2005b; Chou et al., 2004; Fan et al., 2004; Graziano et al., 2006; Huang et al., 2004; Marra et al., 2003; Tan et al., 2005).

We tested the effect of increasing concentrations of DTT up to 10 mM, DMSO up to 10%, and EDTA up to 2 mM and observed no significant effects on enzymatic activity with the exception that EDTA slightly activates at 0.5–1 mM (data not shown). We tested the influence of increasing concentrations of NaCl on SARS-CoV 3CLpro activity up to a concentration of 1 M. We found no significant effect of NaCl up to a concentration of 50–100 mM, but significant inhibition as concentrations moved to values higher than 100 mM. The effect of high concentrations of NaCl on SARS-CoV 3CLpro catalytic activity has also been noted by others (Chou et al., 2004; Graziano et al., 2006; Shi and Song, 2006).

### 4. Conclusions and recommendations

In this study, we evaluated a series of FRET-peptides, including two novel AlexaFluor-QSY FRET peptides, for their ability to serve as substrates for SARS-CoV 3CLpro. We demonstrate for the first time that AlexaFluor dyes and QSY quencher pairs have great potential for use in protease assays. We have shown that N- and C-terminal extensions (tags) to the SARS-CoV 3CLpro enzyme have a significant effect on the activity and quaternary structure of the enzyme. We subjected the enzyme to alterations in salt and organics, while keeping pH constant, and in parallel with other groups we do see alterations in activity in response to added NaCl.

We recommend establishing a standardized assay for examining the activity and structure of SARS-CoV 3CLpro. Although such studies are performed outside the cellular environment and will inherently be artificial, the in vitro assays performed need to be done in a consistent fashion among researchers in order to evaluate results, especially with inhibitors, between groups. We therefore recommend that expression constructs and protocols that produce pure 3CLpro with the authentic N- and C-termini which would be generated from the polypeptide in vivo, be used in kinetic and structural studies of the enzyme. Alternatively, a construct, just as the one used in these studies, that generates 3CLpro with an n-terminal methionine can be used. Any other amino acid additions to SARS-CoV 3CLpro, especially to the N-terminus, are likely to produce significant deviations in kinetic parameters for substrates and inhibitors. We recommend using a pH of 7.5 for optimal enzyme activity, the addition of no more than 100 mM NaCl to the assay buffer, and careful consideration as to the use of reducing agents since they can have a profound effect on certain classes of inhibitors. Reducing agents should be added to the enzyme storage buffer to keep the enzyme active. Small amounts or reducing agents are well tolerated in assays. DMSO can be added up to 10%, and EDTA may be added up to 1 mM to scavenge spurious heavy-metal ions. Use of 0.01–0.1 mg/ml BSA should be considered when
using lower enzyme concentrations in order to help eliminate nonspecific binding of the enzyme to solid surfaces such as the wells in 96-well plates.

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