

All Projects

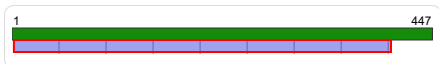
### Untitled Project

Created: today at 01:21

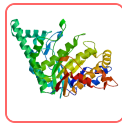
Summary Templates 50 Models 1 Project Data

### Model Results

Order by: GMQE



#### Model 01



Structure Assessment

Compare

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|                    |             |                           |
|--------------------|-------------|---------------------------|
| <b>Oligo-State</b> | <b>GMQE</b> | <b>QMEANDisCo Global:</b> |
| Monomer            | 0.69        | 0.74 ± 0.05               |

**QMEANDisCo Local** ▼  
**QMEAN Z-Scores** ▼

**Template** ▼  
 2pe4.1.A Hyaluronidase-1 Seq Identity  
 43.07%  
 Coverage ▬

#### Model-Template Alignment

```

Mode1_01 L N F R A P P V D P N P F L W A W A P S E F C L G K D D E P L D M
2pe4.1.A  - - F R G P - L P N P P F T T V W W A N T Q W C L E R H G V D V D V
Mode1_01 S L F S E P G S P R I A A T G Q Q V T I P Y V D B L G Y P P Y D S I
2pe4.1.A  S V F D V W A N P G T F R G P D M I I P Y S S L G T P Y V Y - T P
Mode1_01 T G V I Y N G G I P Q W T S L Q D H L D K A K K D I T E Y M P V - D N
2pe4.1.A  T G E P V F G G L P Q N A S L I A H L A R T E Q D I L A A I P A P D F
Mode1_01 L G M A V P D W E E W R P T W A R N W K P K D V Y K A R S I E L V Q Q
2pe4.1.A  S G L A V P D W E A W R P R W A P N W D T K D I Y R Q R S R A L V Q A
Mode1_01 Q N V Q L S L T E A T E K A K Q F F E K A G K D F L V E T I K L Q K L
2pe4.1.A  Q H P D W P A P Q V E A V A Q D Q F G A A R A W M A G T L Q L Q R A
Mode1_01 E R P N H L W G Y L F P D C Y N H H Y K P G Y N G S C F A V Y E I K
2pe4.1.A  E R P R G L W G F Y G F P D C Y N Y D F L S P N Y T G Q C P S G T R A
Mode1_01 R N D L S W L W A S T A L Y P S I Y L N I Q Q S P V A A T L Y V
2pe4.1.A  Q N D L G W L W G S R A L Y P S I Y M P A V L E G T G K S Q M Y V
Mode1_01 R R R V R E A I R V S K L R D A K S P L P W F A V R I V E T Q V L
2pe4.1.A  Q H R V A E A F R V A - V A A G D P N L P Y L P Y Q I - F Y D T T N
Mode1_01 K F L S Q D E L V Y T F G E T V A L G A S G I V I G T E S I M R S M
2pe4.1.A  H F L P L D E L E H S L G E S A A G A A G V V L W V S W E N T R T K
Mode1_01 K S C L L E D N Y M D T I L N P Y I I N V F L A A K M C S Q V L C Q E
2pe4.1.A  E S C Q A I K E Y M D T T L G P F I L N V T S G A L L C S Q A L C S G
Mode1_01 Q G V C I K N W N S S D Y L H L N P D N F A I Q L E - K G G K T V
2pe4.1.A  H G R C V R R T S H P K A L L L N P A S F S I Q L T P G G G P L S L
Mode1_01 R K K P T C E D L E Q F S E K F Y C S C Y S T L S C K E K A D V K D T
2pe4.1.A  R G A L S L E D Q A M A V E F K R C Y - - - - -
Mode1_01 D A V D V C I A D G V C I D A F L K P P M E T E E P Q I F Y
2pe4.1.A  - - - - -
  
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