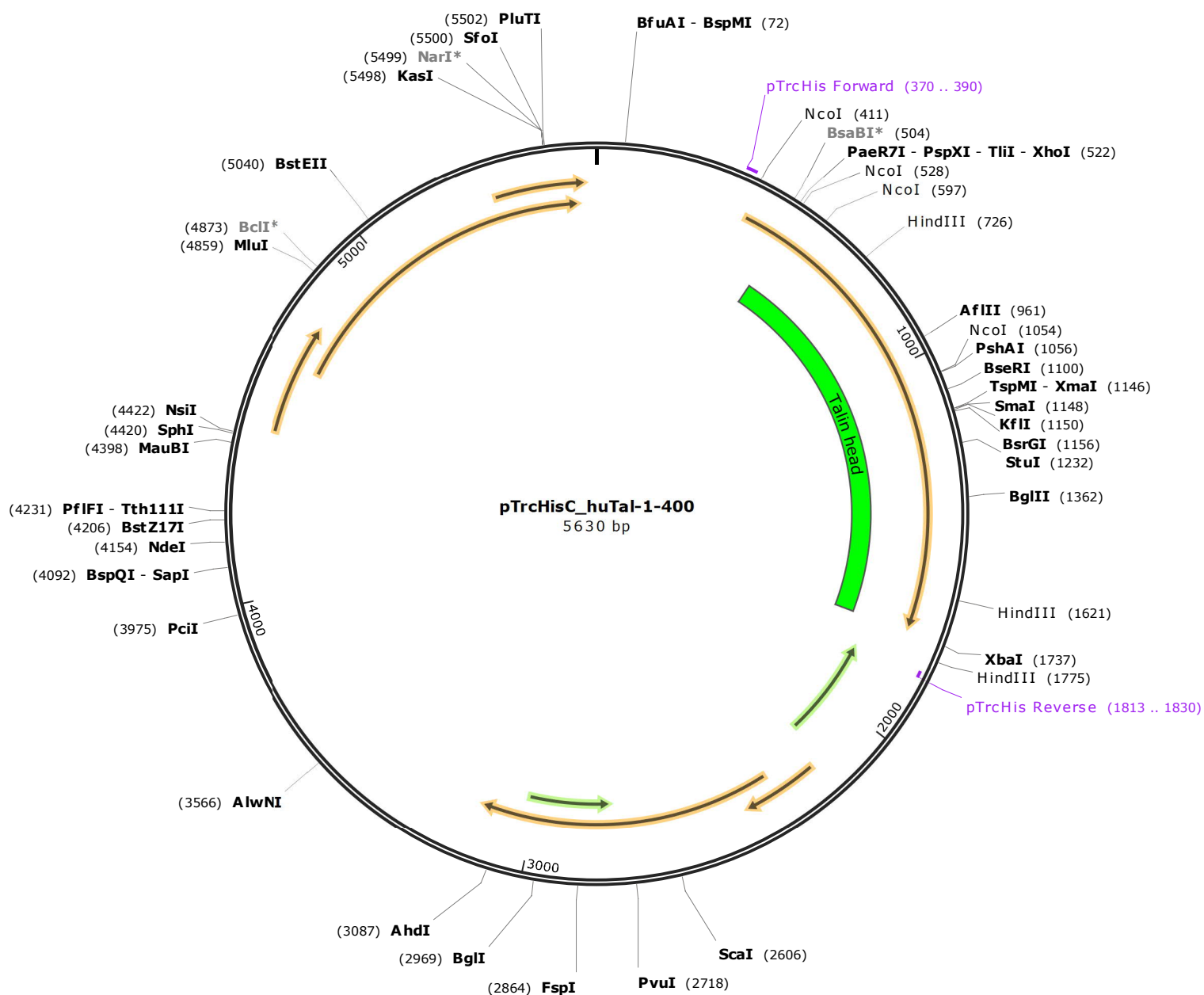
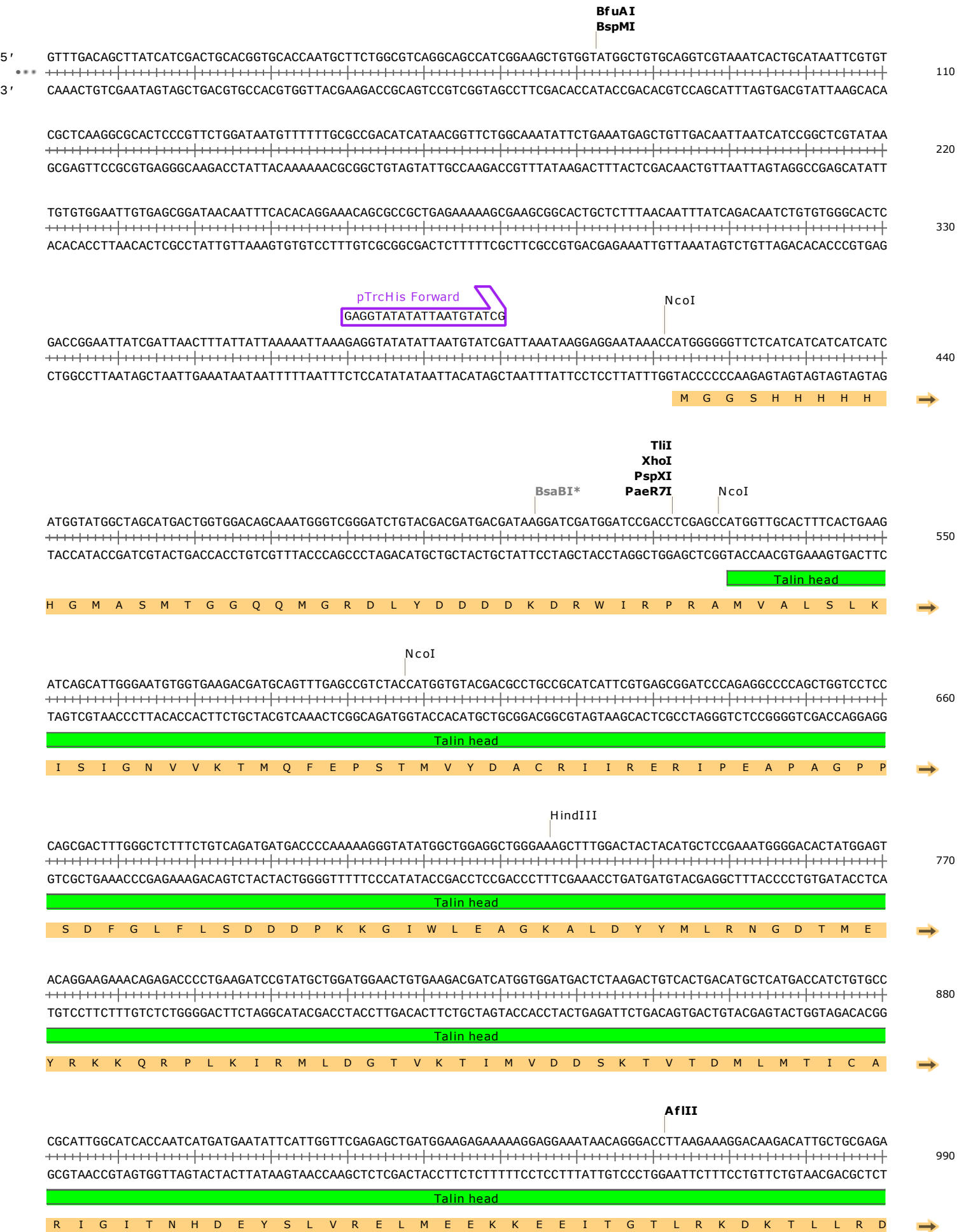


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NcoI PshAI

BseRI

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Talin head

E K K M E K L K Q K L H T D D E L N W L D H G R T L R E Q G V E E H E T

XmaI TspMI SmaI KfII BsrGI

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Talin head

L L L R R K F F Y S D Q N V D S R D P V Q L N L L Y V Q A R D D I L N G S

StuI

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Talin head

H P V S F D K A C E F A G F Q C Q I Q F G P H N E Q K H K A G F L D L K D

BglII

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Talin head

F L P K E Y V K Q K G E R K I F Q A H K N C G Q M S E I E A K V R Y V K

HindIII

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Talin head

M R V D E K T K E V I Q E W N L T N I K R W A A S P K S F T L D F G D Y Q

XbaI

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Talin head

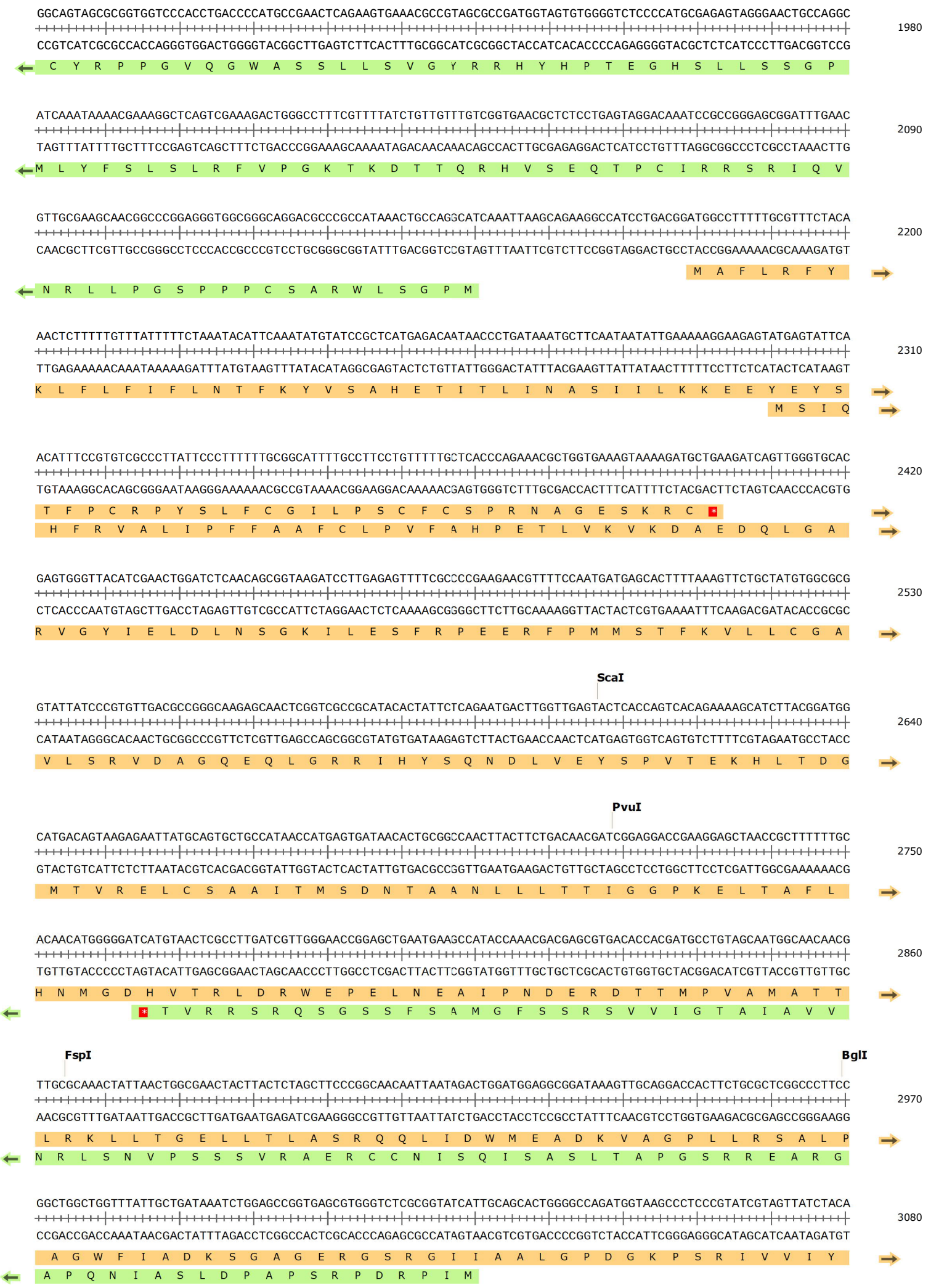
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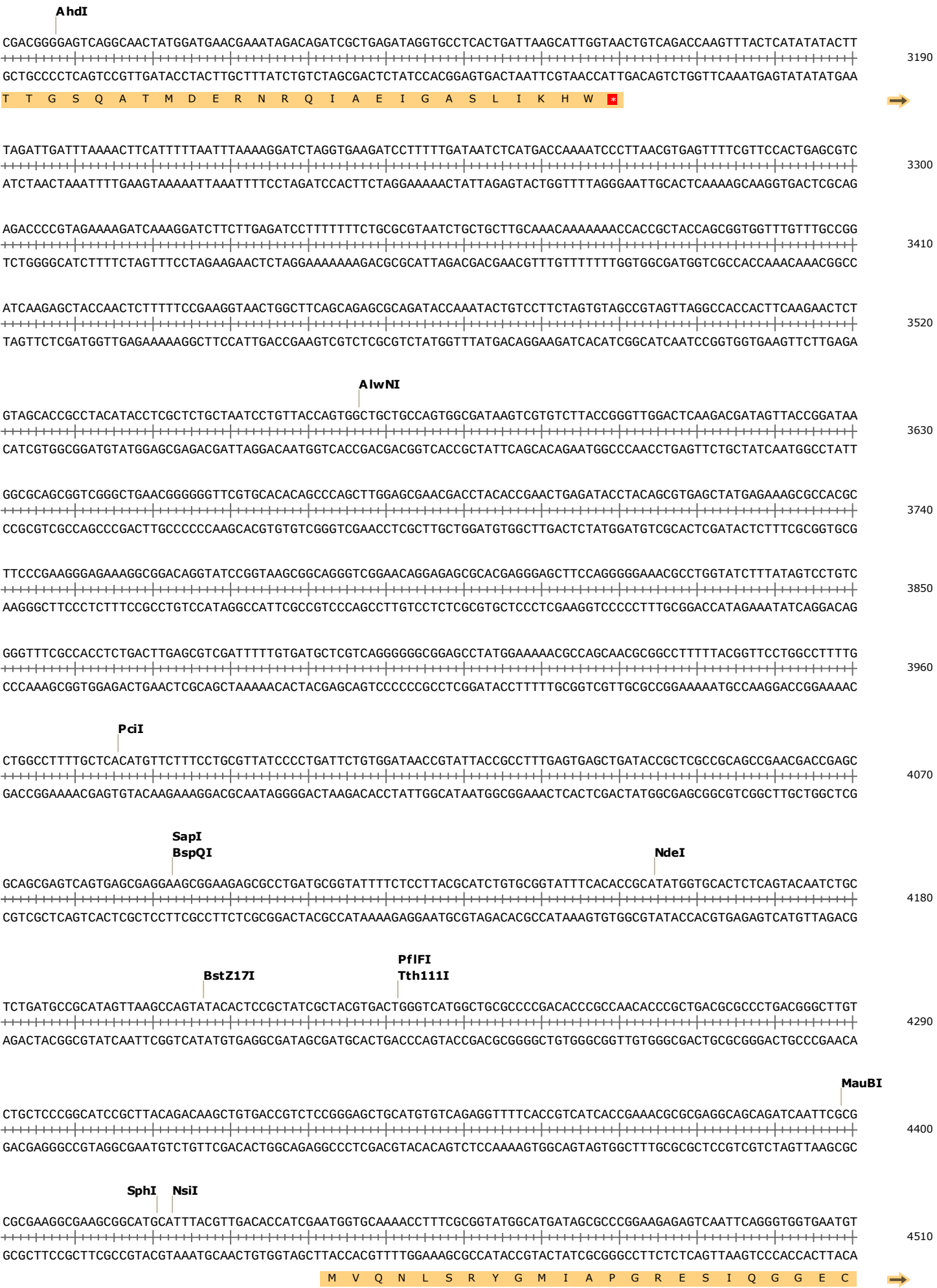
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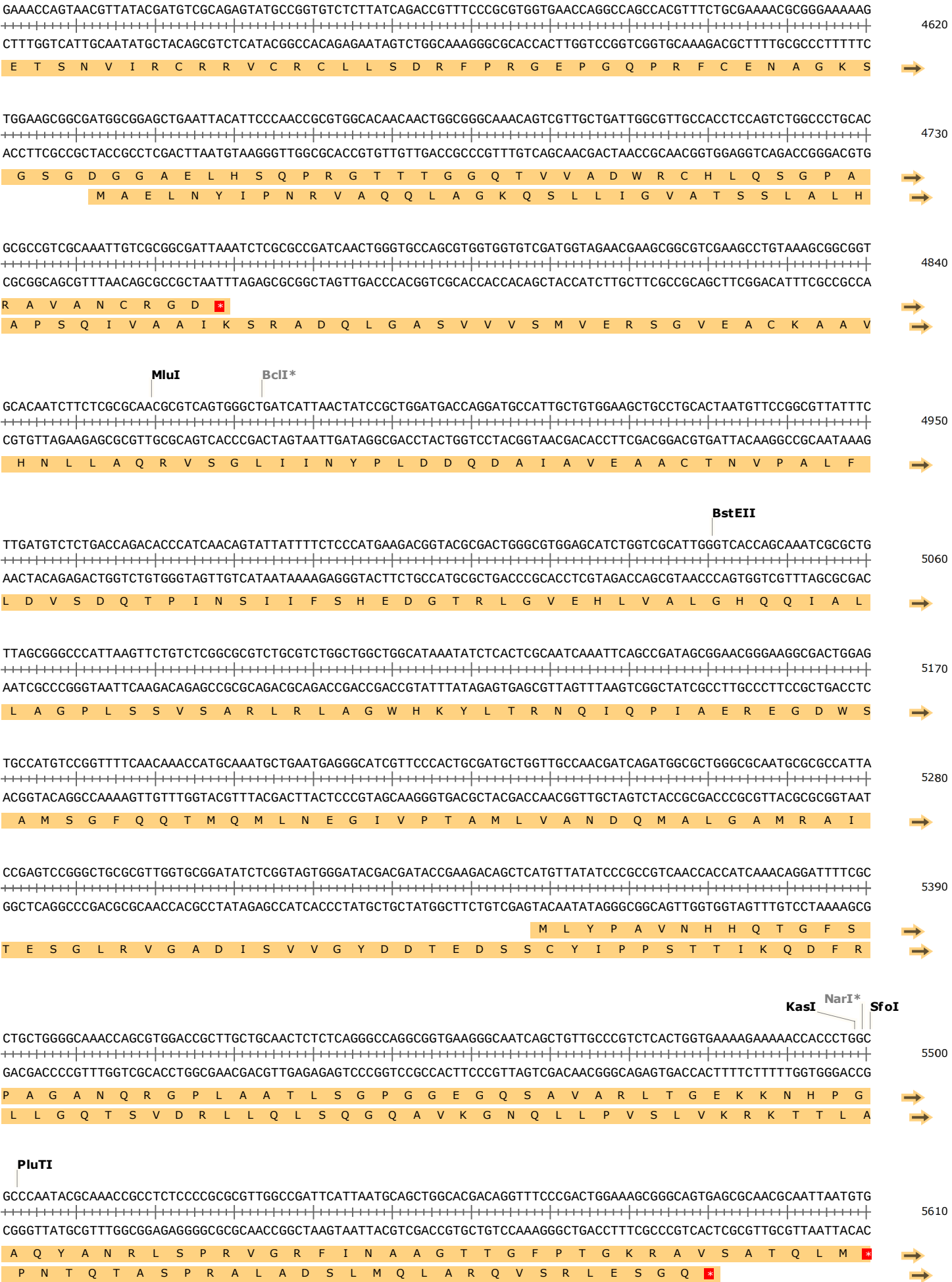
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I L V C F R D S L V S N A Q R

GGACTATGTCTAATTAG
pTrcHis Reverse







AGTTAGCGCGAATTGATCTG 3' 5630
TCAATCGCGCTTAAC TAGAC 5'