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Title	Pathogenesis and therapeutic interventions for ANCA-associated vasculitis.
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Citation	Nature reviews. Rheumatology, 15(2), 91-101 https://doi.org/10.1038/s41584-018-0145-y
Issue Date	2019-02
Doc URL	https://hdl.handle.net/2115/74654
Type	journal article
File Information	Supplementary Table 3.pdf, Supplementary Information



Supplementary Table 3. Other models of AAV

Model	ANCA	Phenotype	Significance	Limitation	Ref.
<i>Drug-induced</i>					
WKY rats orally administered PTU accompanied by intraperitoneal injection of PMA	MPO-ANCA	PGN PH	Useful for determining the involvement of NETs in ANCA production	Mild phenotype	37
BALB/c mice orally administered PTU accompanied by intraperitoneal injection of PMA	MPO-ANCA	No phenotype	Useful for determining the involvement of NETs in ANCA production	No phenotype	54
<i>Molecular mimicry</i>					
WKY rats immunized with FimH	LAMP2-ANCA	NCGN PH	Useful for determining the involvement of molecular mimicry mechanism in the pathogenesis of AAV	“Minor” ANCA	27
<i>Spontaneous</i>					
SCG/Kj mice	MPO-ANCA	NCGN	Useful for therapeutic experiments	Need to differentiate from lupus nephritis	55

PMA, phorbol myristate acetate; PGN, proliferative glomerulonephritis; PH, pulmonary haemorrhage; LAMP2, lysosome-associated membrane protein 2; NCGN, necrotizing and crescentic glomerulonephritis.