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Drug Details

Drug Name(s)	SPIRONOLACTONE
FDA Application No.	(ANDA) 203512
Active Ingredient(s)	SPIRONOLACTONE
Company	ACCORD HLTHCARE INC
Original Approval or Tentative Approval Date	September 20, 2016

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- **Labels are not available**

Products on Application (ANDA) #203512

Click on a column header to re-sort the table:

Drug Name	Active Ingredients	Strength	Dosage Form/Route	Marketing Status	RLD	TE Code
SPIRONOLACTONE	SPIRONOLACTONE	25MG	TABLET;ORAL	Prescription	No	AB
SPIRONOLACTONE	SPIRONOLACTONE	50MG	TABLET;ORAL	Prescription	No	AB
SPIRONOLACTONE	SPIRONOLACTONE	100MG	TABLET;ORAL	Prescription	No	AB

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