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ABBREVIATIONS

μm	micrometer
$^{\circ}\text{C}$	degree celcius
AOR	angle of Repose
BP	British Pharmacopoeia
cm	centimeter
CS	corn starch
CT	chitin
DCP	dibasic calcium phosphate
DI	disintegration time
g	gram
Mg St	magnesium stearate
mg	milligram
ml	millilitre
nm	nanometer
S.D.	Standard Deviation
USP	United State Pharmacopeia