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Review Article

Crystallization and Polymorphism-Scalable Process for Celecoxib and It's Polymorph From-3 (Non- Steroidal Anti-Inflammatory Drug (NSAID)

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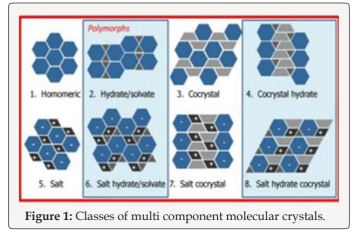
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Abstract

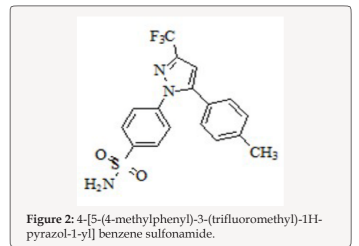
The present process provides an improved process for the preparation of 4-[5-(4-methylphenyl)-3- (trifluoromethyl)-1Hpyrazol-1-yl] benzene sulfonamide (Celecoxib) and its purification and crystallization to produce polymorph. The present process, which describes the manufacturing process of Celecoxib, which is a non- steroidal anti-inflammatory drug (NSAID), has the advantage of scaling up to the industrial level of production. The process uses safe reagents in the process which makes it for industrial scale operations. The yields in the process are high, which makes it a cost-effective process. Formation of isomers are less compared with the all existing process, which makes it effective to make it to the pharmacopoeia grade. Residual solvents play a very important role in the impurity profile of APIs as per the ICH Guidelines ICH Q3C (R4). In this process by carrying out the final step of condensation in the aqueous medium followed by crystallization, the residual solvents limits are well taken care of.

Keywords: Non-steroidal anti-inflammatory drug (NSAID); Celecoxib; Cyclooxygenase 2; X-ray diffraction; Polymorphism; Process

Discussion



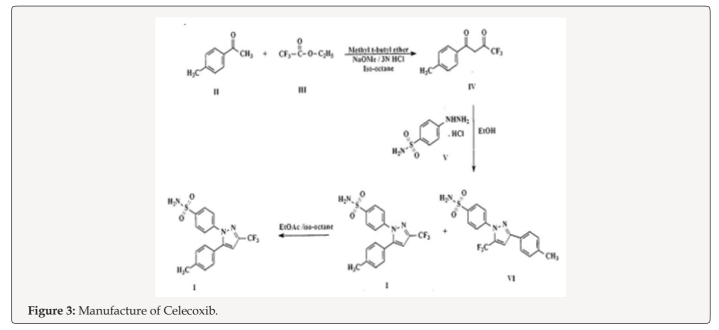
Present process relates to "AN IMPROVED PROCESS FOR THE PREPARATION OF CELECOXIB POLYMORPH FORM". Celecoxib is designated chemically as 4-[5-(4-methylphenyl)-3-(trifluoromethyl)- 1H-pyrazol-1-yl] benzene sulphonamide and is a diaryl-substituted pyrazole [1]. The compound has the following structure (Figure 2).



The drug is currently marketed as Celebrex® in the United States of America by Pharmacia Corporation. Celecoxib is a non-steroidal anti-inflammatory drug (NSAID) [1] mainly used in treatment of arthritis, pain, menstrual cramps, and colonic polyps. Celecoxib blocks the enzyme (cyclooxygenase 2) which

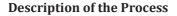
makes prostaglandins, resulting in lowering the concentrations of prostaglandins. As a consequence, reduction in inflammation and its accompanying pain, fever, swelling and tenderness. The manufacture of Celecoxib has been described in various patents and to cite a few references, G. D. Searl & Co. has disclosed method for preparation of Celecoxib [2-3] in US 5,466,823 which is as under: US 5,134,142 [2], US 5,563,165, US 6,150,534, US 5,892,053,

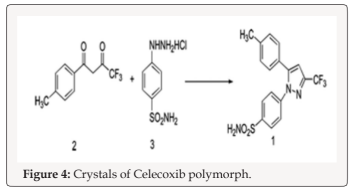
US 2007/0004924, US 2008/0234491, EP 1,528,058, EP 1,167,355, EP 2,246,332, WO 01/42221, WO 03/090730, WO05/014546, WO 06/051340, WO 08/145733, and WO 2010/095024 have also described the synthesis of Celecoxib Reddy et al in their publication in Org. Process Res. Dev., 2009, 13(1), pp 98-101. have disclosed the synthesis (Figure 3).



Detailed Description of the Drawings

Figure 1 describes the powder X-ray diffraction pattern of the Celecoxib Polymorph; Figure 2 illustrates 20values. Figure 3 depicts the DSC thermogram taken at 10:C /min over a temperature range of 30:C to 200:C for Celecoxib polymorphic form.





The present procedure describes the preparation of Celecoxib by a novel process and its crystallization to polymorphic form. The present process for the preparation of Celecoxib by a process involving condensation of 4,4,4-trifluoro-1-[4-(methyl) phenyl]butane-1,3-dione [1] with sulphonamido phenyl hydrazine hydrochloride [2] in an aqueous medium to give Celecoxib [3]. This is followed by crystallization from a mixture of solvents [4-8] containing Aromatic hydrocarbon and aliphatic ketone. In the condensation reaction the reactants are added in water and reactions done at ambient temperature. The crude Celecoxib is isolated by filtration. In the for purification of Celecoxib and its crystallization to polymorphic FORM Preparing a solution of Crude Celecoxib in a solvent mixture comprising of an aliphatic ketone (Acetone) and an aromatic hydrocarbon (Toluene)at reflux temperature followed by cooling crystallization to give crystals of Celecoxib polymorph [8-12] (Figure 4).

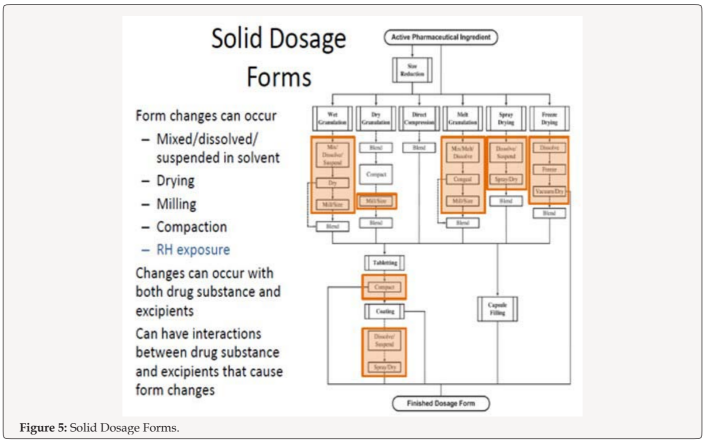
Table 1:

Signal	Assignment/Remarks		
2.3 ppm	3 H, methyl, Singlet		
3.337 and 3.329 ppm	Residual H of DMSO d6, ignore		
7.20 ppm	4 H, Aromatics, multiplet		
7.53 ppm	3 H, Aromatics, 1 H, H-4 (pyrazole)		
8.8 ppm	1 H, Aromatic, doublet		

In this process by carrying out the final step of condensation in the aqueous medium followed by crystallization, the residual solvents limits are well taken care of. The yields in the process are higher compared to the prior art, which makes it a cost-effective process. Formation of isomers are less compared with the prior art, which makes it effective to make it to the pharmacopoeia grade. Residual solvents play a very important role in the impurity profile of APIs as per the ICH Guidelines ICH Q3C (R4). In this process

by carrying out the final step of condensation in the aqueous medium followed by crystallization, the residual solvents limits are well taken care of [13,14]. The crystallization conditions are well established to give crystalline polymorph. The powder X-Ray diffraction pattern of the Celecoxib is given in Figure 1 and 2θ

values are given in Table 1 of Figure 2. The differential scanning calorimeter graph of the Celecoxib polymorph under specific conditions shows the melting point around 1620. The DSC of Celecoxib is given in Figures 3,5, and 6.



Process Induced Transformations				
• Can invo	lve one compone	ent or more		
 Can be s 	olid-solid or solid	l-liquid-solid		
Types of the most important Basic process	phase transitions during processing of pha State of aggregation	Specific process		
Basic process Transformations		Specific process Polymorphic transformations		
Basic process	State of aggregation	Specific process Polymorphic transformations Crystallization of the amorphous form and vice versa Incongruent melting (melting followed by crystallization of a more stable form)		
Basic process Transformations	State of aggregation Solid-solid	Specific process Polymorphic transformations Crystallization of the amorphous form and vice versa Incongruent melting (ineling foilowed by crystallization		
Basic process Transformations (one component) Physical interactions	State of aggregation Solid–solid Solid–liquid–solid	Specific process Polymorphic transformations Crystallization of the amorphous form and vice versa Incongruent melting (ineling followed by crystallization of a more stable form) Solution mechaned polymorphic transformations		
Basic process Transformations (one component) Physical interactions	State of aggregation Solid–solid Solid–tiquid–solid Solid–solid	Specific process Polymorphic transformations Crystallization of the amorphous form and vice versa Incongruent melting (ineling followed by crystallization of a more stable form) Solution mediated polymorphic transformations Eutectic reaction Formation of a molecular compound or a solid solution		

Solid Forms

a) Propensity to produce different forms not significantly different for salts and non-salts.

b) Need more data on co-crystals (Figure 7).

The details of the new methods for preparation of celecoxib are further illustrated in the following examples.

Example 1: Preparation of Celecoxib

In a 20 liter 3-necked flask, equipped with stirrer, thermometer



and reflux condenser, deionized water (7.9 Liter) is charged and mixture of 4,4,4-trifluoro-1-[4-(methyl) phenyl]-butane-1,3-dione (1.6Kg; 6.95×10^3 mmoles) and 4-sulphonamido phenyl hydrazine hydrochloride (1.7Kg; 7.57×10^3 mmoles), a resultant mixture was

heated at 75°C to 80°C and maintained for 5 hours. The reaction mixture was cooled to 25°C to 30°C to give a slurry. The slurry was filtered and washed with water (3.2liter) wet- cake was collected and further processed for purification as given below.

	all compounds [count (%)]	salts [count (%)]	non-salts [count (%)]
multiple forms ^a	220 (89)	86 (91)	116 (91)
multiple crystalline forms ^b	200 (82)	77 (81)	105 (82)
polymorphs ^c	118 (48)	37 (39)	71 (55)
hydrates	94 (38)	46 (48)	38 (30)
solvates	78 (32)	34 (36)	36 (28)
noncrystalline	118 (48)	51 (54)	55 (43)
total compounds	245	95	128

^a Crystalline polymorphs, hydrates, and solvates plus noncrystalline forms. ^b Crystalline polymorphs, hydrates, and solvates. ^c Crystalline polymorphs.

Figure 7: Percentages of forms from Polymorph Screening.

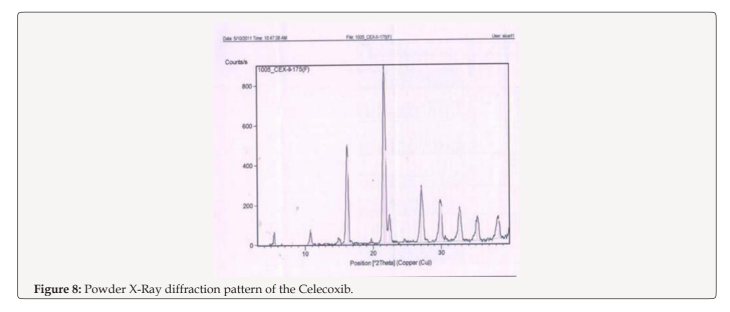
a) Purification and Crystallization to Give Polymorph: Celecoxib wet-cake obtained in the process described above was taken into 20 liter 3-necked flask, equipped with stirrer, thermometer and reflux condenser, mixture of acetone (0.54liter) and toluene (10.8liter) was added and the reaction mixture was heated to 80°C to 85°C for 30 minutes. Activated carbon (0.3Kg) was added and the reaction mixture was further heated to °60 to 85°C. The reaction mixture was cooled to 25°C -30°C. The slurry was filtered, washed with toluene and then dried at 7°C to yield the Celecoxib polymorph compound 1.35 kg (HPLC purity-99.8% & molar yield; 50.9%).

IR: 3340, 3240, 1600, 1500, 1350, 1280, 1235, 1160, 980, 910, 840, 800,760, 635, 560, 530 cm⁻¹(KBr pellet)

Proton NMR: Solvent: DMSO d6, 300 MHz.

Example 2: Preparation of Celecoxib

In a 20liter 3-necked flask, equipped with stirrer, thermometer and reflux condenser, charge deionized water(9Liter) and mixture of 4,4,4-trifluoro-1-[4-(methyl)phenyl]-butane-1,3-dione(1.6Kg; 6.95×10^3 mmoles) and 4-sulphonamido phenyl hydrazine hydrochloride(1.7Kg; 7.57×10³mmoles), a resultant mixture was heated at 90°C to 100°C and maintained for 5 hours. The reaction mixture was cooled to 25°C to 30°C. The slurry was filtered and washed with water (3.2liter) wet-cake was collected and further processed for purification as given below.

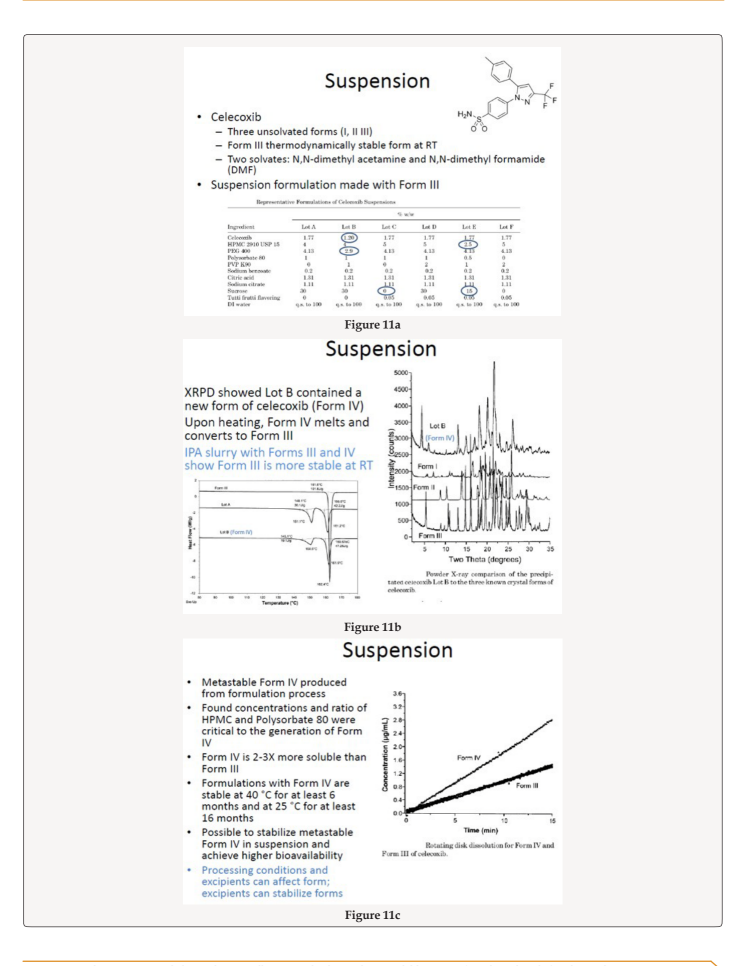


a) Purification and crystallization to give Polymorph: Celecoxib wet-cake obtained in the process described above was taken into 20liter 3-necked flask, equipped with stirrer, thermometer and reflux condenser, mixture of acetone (0.54liter) and toluene (10.8liter) was added and the reaction mixture was heated to 80°C to 85°C for 30 minutes. Activated carbon (0.3Kg) was added and the reaction mixture was further heated to 80°C to 85°C. The reaction mixture was cooled to 25°C -30°C. The separated solid was filtered, washed with toluene and then dried at 70°C to yield the Celecoxib polymorph compound1.24 kg (HPLC purity-99.3% & molar yield; 47%) (Figures 8-11c).

SOPH	ISTICATED INSTR	MOTA BAZAR -					
No.	Visible	Dataset Name	Start pos. [*2Th.]		Step [*2Th.]	Measured Date/Time	
1	TRUE	1005_CEX-8- 175(F)	3.01	39.99	0.02	5/10/2011 10:28	
No.	Pos. [*2Th.]	d-spacing	Significance	Rel. Int. [%]	Height	FWHM (*2Th.)	
1	5,4088	16.33928	3.1387	9.86	64.67	0.1968	
2	10.8034	8.18943	and the second se	7.64	71.62	0.1378	
3		6.80362	1.891	2	5.45	0.4723	
4	and the second se	5.90145	1.6412	7.38	24.2	0.3936	
5	16.2935	5.44029	9.2472	100	437.32	0.2957	
6	and the second design of the s	4.50894	0.9851	2.04	22.3	0.118)	
7	and the second se	4.1137	1.0274	80.09	875.58	0.1181	
8	21.8734	4.06345	0.9792	42.77	457.65	and the second se	
9	22.4507	3.96026	1.4827	21.45	140.72	and the second se	
30	23.824	3.735	0.8233	0.45	4.88		
11	24.5936	3.61984	0.833	4.77	13.03	And and a subscription of the local division	
17	27.054	3.29597	0.8743		and the second se		
13	29.7841	2.99977	1.0088				
14	30.0469	2.97413	1.1780	19.03	And and a second s	and the second se	
15	32.6341	2.74401	1.6601	and the second s	And and a state of the local division of the	and the second se	
10	35.3025	2.54245	1.6827	and the second design of the s	and the second designed in the second designed in the second designed and the		
17	38,4104	2.34361	4.8435	40.55	9	0.55	

Figure 9: 2θ values.

tole Weight	D:2011Way/0905-CEX-II-171-F- SICART Celecox/b(CEX/I/(171(F) 2.393 mg Amoli Organics Pvt.Ltd. ID No-0720/11-12	1112.dod				
0		Sophisticated Instrumen	tation Centre for Applied Research and T	esting (SICART)		
				Peak = 163.517 °C Peak Height = 21.090	Wm 0	
5				Area = 232.572 mJ Deita H = 97.188 J/g		
0				-		
5						
D						
5						
00						
2 55 60	80	100	120 Temperature (°C)	160	180	20
					5/9/114.5	2:53 PM
Hold for 1.0	nin at 50.00°C		 Heat from 50.00°C 1 	o 200.00°C at 10.00°C/min		



Conclusion

The distinct advantage of the present method of preparation over the prior art can be summarized as per below:

The present process, which describes the manufacturing process of Celecoxib, which is a non- steroidal anti-inflammatory drug (NSAID), has the advantage of scaling up to the industrial level of production. The process uses safe reagents in the process which makes it for industrial scale operations. The present process provides an improved process for the preparation of 4-[5-(4-methylphenyl)-3-(trifluoromethyl)-1H-pyrazol-1yl] benzene sulfonamide (Celecoxib) and its purification and crystallization to produce polymorph. The yields in the process are high compared to existing process which makes it a cost-effective process. Formation of isomers are less compared with the prior art, which makes it effective to make it to the pharmacopoeia grade. In this process by carrying out the final step of condensation in the aqueous medium followed by crystallization, the residual solvents limits are well taken care of. The yields in the process are higher compared to the prior art, which makes it a cost-effective process. Residual solvents play a very important role in the impurity profile of APIs as per the ICH Guidelines ICH Q3C (R4).

References

- 1. Pouton CW (2006) Formulation of poorly water-soluble drugs for oral administration: physicochemical and physiological issues and the lipid formulation classification system. Eur J Pharm Sci 29(3-4): 278-287.
- 2. Matsuo (2012) US 5,134,142, US 5,563,165, US 6,150,534, US5,892,053, US 2007/0004924.



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- 3. G Sievers R (Eds.); Data Commission of the International Union of Crystallography, Bonn pp. 171-200.
- 4. Anonymous (2001) 50 top selling drugs ranked by worldwide sales for 2000, Patent scan-a monthly proactive patents bulletin for pharmaceutical industry. Med Ad News 2(12).
- Behme RJ, Brooke D, Farney RF, Kensler TT (1985) Characterization of polymorphism of gepirone hydrochloride. J Pharm Sci 74(10): 1041-1046.
- 6. Brittain HG (1997) Spectral methods for the characterization of polymorphs and solvates. J Pharm Sci 86(4): 405-412.
- 7. Brittain H, Fiese E (1999) Effects of pharmaceutical processing on drug polymorphs and solvates. In: Swarbrick J (Eds.); Polymorphism in Pharmaceutical Solids. Marcel Dekker, New York, pp. 331-365.
- Brittain HG and Grant DJW (1999) Effects of polymorphism and solidstate solvation on solubility and dissolution rate. In: Swarbrick J (Eds.); Polymorphism in Pharmaceutical Solids. Marcel Dekker, New York pp. 279-330.
- 9. Giron D (1995) Thermal analysis and calorimetric methods in the characterization of polymorphs and solvates. Thermochim Acta 248: 1-59.
- 10. Halgren T (1990) MOMEC force field parameters. J Am Chem Soc 112: 4710-4725.
- 11. Hancock B, Zografi G (1997) Characterization and significance of the amorphous state in pharmaceutical systems. J Pharm Sci 86(1):1-12.
- 12. Jozwiakowski M (2000) Alteration of the solid state of the drug substance: polymorphs, solvates, and amorphous forms. In: Liu R (Eds.); Water Insoluble Drug Formulation. Interpharm Press, Colorado pp. 525-562.
- 13. Kakumanu V, Bansal A (2002) Enthalpy relaxation studies of celecoxib amorphous mixtures. Pharm Res 19(12): 1873-1878.
- 14. Leonard F, Patricia M (1999) Polymorphic crystalline forms of celecoxib.



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