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(54) **New forms of 5-azacytidine**
Neue Form von 5-azacytidine
Nouvelles formes de la 5-azacytidine

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Remarks:

The file contains technical information submitted after the application was filed and not included in this specification

Description

FIELD OF THE INVENTION

5 **[0001]** The invention relates to a pharmaceutical composition comprising a crystalline form of 5-azacytidine for use in a method of treating myelodysplastic syndromes, wherein the crystalline form of 5-azacytidine is characterized by peaks observed using X-ray powder diffraction (Cu K α radiation), comprising peaks having the following 2 θ angles:

	2θ Angle (°)
10	6.566
	11.983
	13.089
15	15.138
	17.446
	20.762
20	21.049
	22.776
	24.363
	25.743
25	26.305
	28.741
	31.393
30	32.806
	33.043
	33.536
	36.371
35	39.157
	41.643

, and wherein the method comprises the oral administration of the pharmaceutical composition.

40 **[0002]** In addition, the invention relates to a pharmaceutical composition comprising a mixed phase of forms of 5-azacytidine for use in a method of treating myelodysplastic syndromes, wherein the mixed phase of forms of 5-azacytidine is characterized by peaks observed using X-ray powder diffraction (Cu K α radiation), having the following 2 θ angles:

	2θ Angle (°)
45	12.244
	13.082
	13.458
50	14.452
	16.521
	17.648
55	18.677
	19.093
	20.231

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(continued)

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2 θ Angle (°)
21.353
22.309
23.070
23.909
26.641
26.813
27.158
29.309
29.609
30.384
32.074

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, and wherein the method comprises the oral administration of the pharmaceutical composition.

[0003] The invention is also directed to a pharmaceutical composition comprising a crystalline form of 5-azacytidine for use in a method of treating myelodysplastic syndromes, wherein the crystalline form is characterized by peaks observed using X-ray powder diffraction (Cu K α radiation), comprising peaks having the following 2 θ angles:

25

2 θ Angle (°)
12.533
12.963
13.801
18.929
20.920
21.108
21.527
22.623
22.970
24.054
26.668
27.210
28.519
29.548
30.458
33.810
35.079
37.528

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, and wherein the method comprises the oral administration of the pharmaceutical composition.

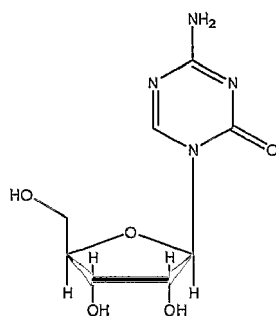
[0004] The invention further relates to a pharmaceutical composition comprising an amorphous solid of 5-azacytidine for use in a method of treating myelodysplastic syndromes, wherein the method comprises the oral administration of the pharmaceutical composition.

[0005] Described herein is the isolation of crystalline polymorphic and pseudopolymorphic forms of 5-azacytidine (also known as azacitidine and 4-amino-1- β -D-ribofuranosyl-S-triazin-2(1*H*)-one). 5-azacytidine may be used in the treatment of disease, including the treatment of myelodysplastic syndromes (MDS).

5 BACKGROUND OF THE INVENTION

[0006] Polymorphs exist as two or more crystalline phases that have different arrangements and/or different conformations of the molecule in a crystal lattice. When a solvent molecule(s) is contained within the crystal lattice the resulting crystal is called a pseudopolymorph, or solvate. If the solvent molecule(s) within the crystal structure is a water molecule, then the pseudopolymorph/solvate is called a hydrate. The polymorphic and pseudopolymorphic solids display different physical properties, including those due to packing, and various thermodynamic, spectroscopic, interfacial and mechanical properties (See H. Brittain, *Polymorphism in Pharmaceutical Solids*, Marcel Dekker, New York, NY, 1999, pp. 1-2). Polymorphic and pseudopolymorphic forms of the drug substance (also known as the "active pharmaceutical ingredient" (API)), as administered by itself or formulated as a drug product (also known as the final or finished dosage form, or as the pharmaceutical composition) are well known and may affect, for example, the solubility, stability, flowability, fractability, and compressibility of drug substances and the safety and efficacy of drug products, (see, e.g., Knapman, *K Modern Drug Discoveries*, March 2000: 53).

[0007] 5-azacytidine (also known as azacitidine and 4-amino-1- β -D-ribofuranosyl-1,3,5-triazin-2(1*H*)-one; Nation Service Center designation NSC-102816; CAS Registry Number 320-67-2) has undergone NCI-sponsored clinical trials for the treatment of myelodysplastic syndromes (MDS). See Kornblith et al., *J. Clin. Oncol.* 20(10): 2441-2452 (2002) Gryn et al., *Leukemia Research* 26(2002) 893-897 and Silverman et al., *J. Clin. Oncol.* 20(10): 2429-2440 (2002). 5-azacytidine may be defined as having a formula of $C_8H_{12}N_4O_5$, a molecular weight of 244.20 and a structure of:



[0008] The polymorphic form of 5-azacytidine drug substance and drug product has never been characterized. It is an object of the present invention to characterize the polymorphic forms of 5-azacytidine.

40 SUMMARY OF THE INVENTION

[0009] The invention relates to a pharmaceutical composition comprising a crystalline form of 5-azacytidine for use in a method of treating myelodysplastic syndromes, wherein the crystalline form of 5-azacytidine is characterized by peaks observed using X-ray powder diffraction (Cu $K\alpha$ radiation), comprising peaks having the following 2θ angles:

2θ Angle ($^\circ$)
6.566
11.983
13.089
15.138
17.446
20.762
21.049
22.776

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(continued)

5
10
15
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2θ Angle (°)
24.363
25.743
26.305
28.741
31.393
32.806
33.043
33.536
36.371
39.157
41.643

, and wherein the method comprises the oral administration of the pharmaceutical composition.

[0010] In addition, the invention relates to a pharmaceutical composition comprising a mixed phase of forms of 5-azacytidine for use in a method of treating myelodysplastic syndromes, wherein the mixed phase of forms of 5-azacytidine is characterized by peaks observed using X-ray powder diffraction (Cu K α radiation), having the following 2 θ angles:

25
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35
40
45
50
55

2θ Angle (°)
12.244
13.082
13.458
14.452
16.521
17.648
18.677
19.093
20.231
21.353
22.309
23.070
23.909
26.641
26.813
27.158
29.309
29.609
30.384
32.074

, and wherein the method comprises the oral administration of the pharmaceutical composition.

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[0011] The invention is also directed to a pharmaceutical composition comprising a crystalline form of 5-azacytidine for use in a method of treating myelodysplastic syndromes, wherein the crystalline form is characterized by peaks observed using X-ray powder diffraction (Cu K α radiation), comprising peaks having the following 2 θ angles:

2 θ Angle (°)
12.533
12.963
13.801
18.929
20.920
21.108
21.527
22.623
22.970
24.054
26.668
27.210
28.519
29.548
30.458
33.810
35.079
37.528

, and wherein the method comprises the oral administration of the pharmaceutical composition.

[0012] The invention further relates to a pharmaceutical composition comprising an amorphous solid of 5-azacytidine for use in a method of treating myelodysplastic syndromes, wherein the method comprises the oral administration of the pharmaceutical composition.

[0013] It has been unexpectedly found that 5-azacytidine exists in at least eight different polymorphic and pseudopolymorphic crystalline forms (Forms I-VIII), in addition to an amorphous form. Form I is a polymorph found in prior art retained samples of 5-azacytidine drug substance. Form II is a polymorph found in some prior art retained samples of the 5-azacytidine drug substance; in those samples, Form II is always found in mixed phase with Form I. Form III is a hydrate, and is formed when prior art retained and current samples of the drug product are reconstituted with water to form a "slurry" prior to administration to the patient. Form VI is found in prior art retained samples of the 5-azacytidine drug product, either substantially free of other polymorphs, or in mixed phase with Form I.

[0014] The description provides novel crystalline forms referred to as Form IV, Form V, Form VII and Form VIII. Forms I-VIII each have characteristic X-ray powder diffraction (XRPD) patterns and are easily distinguished from one another using XRPD.

[0015] Also included in the description are methods for robustly and reproducibly synthesizing 5-azacytidine drug substance substantially as Form IV, Form V, or Form VIII. Also provided are methods for robustly and reproducibly synthesizing a Form I/VII mixed phase. The invention also provides pharmaceutical compositions comprising the various forms of 5-azacytidine together with one or more pharmaceutically acceptable excipients, diluents, or carriers.

BRIEF DESCRIPTION OF THE FIGURES

[0016]

Figure 1 presents the X-Ray Powder Diffraction (XRPD) pattern of 5-azacytidine, Form I, labeled with the most prominent 2 θ angles (Cu K α radiation).

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Figure 2 presents the XRPD pattern of 5-azacytidine, mixed phase Form I and Form II, labeled with the most prominent 2θ angles (Cu $K\alpha$ radiation).

Figure 3 presents the XRPD pattern of 5-azacytidine, Form III, labeled with the most prominent 2θ angles (Cu $K\alpha$ radiation).

Figure 4 presents the XRPD pattern of 5-azacytidine, Form IV, labeled with the most prominent 2θ angles (Cu $K\alpha$ radiation).

Figure 5 presents the XRPD pattern of 5-azacytidine, Form V, labeled with the most prominent 2θ angles (Cu $K\alpha$ radiation).

Figure 6 presents the XRPD pattern of 5-azacytidine, Form VI, labeled with the most prominent 2θ angles (Cu $K\alpha$ radiation).

Figure 7 presents the XRPD pattern of 5-azacytidine, mixed phase Form I and Form VII, labeled with the most prominent 2θ angles (Cu $K\alpha$ radiation).

Figure 8 presents the XRPD pattern of 5-azacytidine, Form VIII, labeled with the most prominent 2θ angles (Cu $K\alpha$ radiation).

DETAILED DESCRIPTION

[0017] The invention relates to a pharmaceutical composition comprising a crystalline form of 5-azacytidine for use in a method of treating myelodysplastic syndromes, wherein the crystalline form of 5-azacytidine is characterized by peaks observed using X-ray powder diffraction (Cu $K\alpha$ radiation), comprising peaks having the following 2θ angles:

2θ Angle (°)
6.566
11.983
13.089
15.138
17.446
20.762
21.049
22.776
24.363
25.743
26.305
28.741
31.393
32.806
33.043
33.536
36.371
39.157
41.643

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, and wherein the method comprises the oral administration of the pharmaceutical composition.

[0018] It also relates to the pharmaceutical composition, wherein the crystalline form of 5-azacytidine is characterized by an X-ray powder diffraction (Cu K α radiation) pattern substantially in accordance with Figure 3; optionally wherein the crystalline form of 5-azacytidine is characterized by peaks observed using X-ray powder diffraction (Cu K α radiation), comprising peaks having the following 2 θ angles, d-spacings and relative intensities:

2 θ Angle (°)	d-spacing (Å)	Relative Intensity
6.566	13.450	32.9
11.983	7.380	52.5
13.089	6.758	71.0
15.138	5.848	38.9
17.446	5.079	48.2
20.762	4.275	10.8
21.049	4.147	34.8
22.776	3.901	89.5
24.363	3.651	13.7
25.743	3.458	22.8
26.305	3.385	39.9
28.741	3.104	100.0
31.393	2.847	22.5
32.806	2.728	11.8
33.043	2.709	10.1
33.536	2.670	15.1
36.371	2.468	11.0
39.157	2.299	19.3
41.643	2.167	12.1

[0019] One embodiment of the invention is the aforementioned pharmaceutical composition, wherein the crystalline form of 5-azacytidine is a monohydrate prepared according to a method comprising:

contacting a crystalline form of 5-azacytidine with water; and
isolating the crystalline form of 5-azacytidine; optionally wherein the crystalline form of 5-azacytidine used in the preparation of the crystalline form 5-azacytidine according to claims 1 to 3 is Form I, Form II, Form IV, Form V, Form VI, Form VII, or Form VIII of 5-azacytidine; or a mixture of two or more thereof.

[0020] In addition, the invention relates to a pharmaceutical composition comprising a mixed phase of forms of 5-azacytidine for use in a method of treating myelodysplastic syndromes, wherein the mixed phase of forms of 5-azacytidine is characterized by peaks observed using X-ray powder diffraction (Cu K α radiation), having the following 2 θ angles:

2 θ Angle (°)
12.244
13.082
13.458
14.452
16.521

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(continued)

5
10
15
20
25

2θ Angle (°)
17.648
18.677
19.093
20.231
21.353
22.309
23.070
23.909
26.641
26.813
27.158
29.309
29.609
30.384
32.074

, and wherein the method comprises the oral administration of the pharmaceutical composition.

[0021] The invention also relates to the pharmaceutical composition, wherein the mixed phase of forms of 5-azacytidine is characterized by an X-ray powder diffraction pattern substantially in accordance with Figure 2; optionally wherein the mixed phase of forms of 5-azacytidine is characterized by peaks observed using X-ray powder diffraction (Cu K α radiation), comprising peaks having the following 2 θ angles, d-spacings and relative intensities:

35
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45
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2θ Angle (°)	d-spacing (Å)	Relative Intensity
12.244	7.223	34.8
13.082	6.762	37.0
13.458	6.574	29.2
14.452	6.124	25.4
16.521	5.361	19.0
17.648	5.022	12.1
18.677	4.747	12.7
19.093	4.645	41.3
20.231	4.386	42.1
21.353	4.158	15.5
22.309	3.982	35.1
23.070	3.852	100.0
23.909	3.719	18.9
26.641	3.343	18.2
26.813	3.322	12.6
27.158	3.281	46.0
29.309	3.045	27.3

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(continued)

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2θ Angle (°)	d-spacing (Å)	Relative Intensity
29.609	3.015	12.7
30.384	2.939	10.5
32.074	2.788	12.0

10

[0022] The invention is also directed to a pharmaceutical composition comprising a crystalline form of 5-azacytidine for use in a method of treating myelodysplastic syndromes, wherein the crystalline form is characterized by peaks observed using X-ray powder diffraction (Cu K α radiation), comprising peaks having the following 2θ angles:

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2θ Angle (°)
12.533
12.963
13.801
18.929
20.920
21.108
21.527
22.623
22.970
24.054
26.668
27.210
28.519
29.548
30.458
33.810
35.079
37.528

, and wherein the method comprises the oral administration of the pharmaceutical composition.

45

[0023] The invention is also directed to the pharmaceutical composition, wherein the crystalline form of 5-azacytidine is characterized by an X-ray powder diffraction (Cu K α radiation) pattern substantially in accordance with Figure 6; optionally wherein the crystalline form of 5-azacytidine is characterized by peaks observed using X-ray powder diffraction (Cu K α radiation), comprising peaks having the following 2θ angles, d-spacings and relative intensities:

50

55

2θ Angle (°)	d-spacing (Å)	Relative Intensity
12.533	7.057	10.1
12.963	6.824	10.2
13.801	6.411	100.0
18.929	4.6843	10.0
20.920	4.243	34.2
21.108	4.205	49.4

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(continued)

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2θ Angle (°)	d-spacing (Å)	Relative Intensity
21.527	4.125	47.0
22.623	3.922	10.7
22.970	3.869	13.8
24.054	3.697	77.8
26.668	3.340	23.0
27.210	3.275	33.7
28.519	3.127	12.9
29.548	3.021	27.2
30.458	2.932	50.3
33.810	2.649	11.6
35.079	2.556	12.6
37.528	2.411	24.7

[0024] The invention also relates to the pharmaceutical composition, wherein the crystalline form of 5-azacytidine is prepared according to a method comprising:

25 lyophilizing a solution of 5-azacytidine and mannitol; and
isolating the lyophilized solid; optionally wherein the solution of 5-azacytidine and mannitol comprises 5-azacytidine and mannitol in a ratio of about 1:1 by weight.

30 **[0025]** In addition thereto, the invention further relates to a pharmaceutical composition comprising an amorphous solid of 5-azacytidine for use in a method of treating myelodysplastic syndromes, wherein the method comprises the oral administration of the pharmaceutical composition.

35 **[0026]** It also relates to this pharmaceutical composition, wherein the amorphous solid of 5-azacytidine is prepared by a method comprising:
adding a crystalline form of 5-azacytidine, being characterized by peaks observed using X-ray powder diffraction (Cu K α radiation) having the following 2 θ angles:

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45
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2θ Angle (°)
12.182
13.024
14.399
16.470
18.627
19.049
20.182
21.329
23.033
23.872
26.863
27.135
29.277

(continued)

2 θ Angle (°)
29.591
30.369
32.072

, to a solvent selected from the group consisting of propylene glycol, polyethylene glycol, and DMSO; allowing equilibration to occur; and recovering 5-azacytidine therefrom.

[0027] One embodiment of the invention is directed to each of the before mentioned pharmaceutical compositions, wherein the pharmaceutical composition further comprises a pharmaceutically acceptable excipient, diluent, or carrier; or wherein the pharmaceutical composition is made in an unit dosage form; optionally wherein each dosage unit contains from about 5 mg to about 200 mg of the 5-azacytidine form, preferably about 100 mg of the 5-azacytidine form.

5-azacytidine Crystalline Forms I-VIII

[0028] It has been discovered that 5-azacytidine exists in at least eight different polymorphic and pseudopolymorphic crystalline forms, and also in an amorphous form.

Form I

[0029] A single sample of the 5-azacytidine drug substance was synthesized from 5-azacytosine and 1,2,3,5,-Tetra-O-acetyl- β -D-ribofuranose according to the prior art method provided in Example 1. The last step of this method is a recrystallization of the crude synthesis product from a DMSO/methanol co-solvent system. Specifically, the crude synthesis product is dissolved in DMSO (preheated to about 90°C), and then methanol is added to the DMSO solution. The co-solvent mixture is equilibrated at approximately -20°C to allow 5-azacytidine crystal formation. The product is collected by vacuum filtration and allowed to air dry.

[0030] The X-Ray Powder Diffraction (XRPD; see Example 5) pattern of the resulting 5-azacytidine is shown in Figure 1 along with some of the 2 θ values. Table 1 provides the most prominent 2 θ angles, d-spacing and relative intensities for this material, which is designated as Form I.

Table 1: 5-azacytidine Form I -the most prominent 2 θ angles, d-spacing and relative intensities (Cu K α radiation)

2 θ Angle (°)	d-spacing (Å)	Relative Intensity
12.182	7.260	39.1
13.024	6.792	44.1
14.399	6.146	31.5
16.470	5.378	27.1
18.627	4.760	16.0
19.049	4.655	35.9
20.182	4.396	37.0
21.329	4.162	12.4
23.033	3.858	100.0
23.872	3.724	28.0
26.863	3.316	10.8
27.135	3.284	51.5
29.277	3.048	25.6
29.591	3.016	11.5
30.369	2.941	10.8

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(continued)

2θ Angle (°)	d-spacing (Å)	Relative Intensity
32.072	2.788	13.4

[0031] Thermal analysis of Form I indicates that this form of 5-azacytidine is anhydrous. See Example 6.

Form II

[0032] Retained samples of the drug substance previously used to formulate the drug product in the NCI-sponsored Cancer and Leukaemia Group B (CALGB) investigations (Phase 2 trial 8291 and Phase 3 trial 9221) for the treatment of MDS (Investigational New Drug (IND) 7574) were also analyzed by XRPD. The retained drug substance samples comprised either Form I, or a mixed phase of Form I and another polymorph: Form II. See Example 5.

[0033] The XRPD powder pattern of mixed phase Forms I and II is shown in Figure 2 along with some of the 2θ values. Peaks distinctive to Form II are observed at 13.5, 17.6 and 22.3 ° 2θ.

Table 2 provides the most prominent 2θ angles, d-spacing and relative intensities for this mixed phase.

Table 2: 5-azacytidine, Mixed Phase Forms I and II the most prominent 2θ angles, d-spacing and relative intensities (Cu Kα radiation)

2θ Angle (°)	d-spacing (Å)	Relative Intensity
12.244	7.223	34.8
13.082	6.762	37.0
13.458*	6.574	29.2
14.452	6.124	25.4
16.521	5.361	19.0
17.648*	5.022	12.1
18.677	4.747	12.7
19.093	4.645	41.3
20.231	4.386	42.1
21.353	4.158	15.5
22.309*	3.982	35.1
23.070	3.852	100.0
23.909	3.719	18.9
26.641	3.343	18.2
26.813	3.322	12.6
27.158	3.281	46.0
29.309	3.045	27.3
29.609	3.015	12.7
30.384	2.939	10.5
32.074	2.788	12.0

[0034] These results indicate that the prior art 5-azacytidine synthesis procedures for the drug substance produce either Form I substantially free of other forms, or a Form I/II mixed phase *i.e.* a solid material in which 5-azacytidine is present in a mixed phase of both Form I and Form II.

[0035] Thermal analysis of mixed phase Form I/II is presented in Example 6.

Form III

[0036] An additional crystalline form of 5-azacytidine, designated Form III, is found in slurries of 5-azacytidine. See Example 8. Moreover, it has been found that all forms of 5-azacytidine (including the 5-azacytidine in the prior art drug product) convert to Form III in water. See Example 8. Thus, reconstitution of the drug product used in the aforementioned NCI trials would have led to the formation of a saturated solution (or "slurry") in which the remaining solid 5-azacytidine was Form III. The XRPD powder pattern of Form III is shown in Figure 3 along with some of the 2θ values. Table 3 provides the most prominent 2θ angles, d-spacing and relative intensities for this crystalline material. The XRPD powder pattern for Form III is distinctly different from that of all of the other forms of 5-azacytidine.

Table 3: 5-azacytidine, Form III - the most prominent 2θ angles, d-spacing and relative intensities (Cu $K\alpha$ radiation)

2θ Angle (°)	d-spacing (Å)	Relative Intensity
6.566	13.450	32.9
11.983	7.380	52.5
13.089	6.758	71.0
15.138	5.848	38.9
17.446	5.079	48.2
20.762	4.275	10.8
21.049	4.147	34.8
22.776	3.901	89.5
24.363	3.651	13.7
25.743	3.458	22.8
26.305	3.385	39.9
28.741	3.104	100.0
31.393	2.847	22.5
32.806	2.728	11.8
33.043	2.709	10.1
33.536	2.670	15.1
36.371	2.468	11.0
39.157	2.299	19.3
41.643	2.167	12.1

[0037] Thermal analysis and proton (^1H) NMR spectroscopy indicate that Form III is a pseudopolymorphic form of 5-azacytidine, specifically a monohydrate. See Examples 6-7.

Form IV

[0038] Form IV is a crystalline form of 5-azacytidine. Form IV was recovered by slow recrystallization from a DMSO/toluene co-solvent system (see Example 2) or by fast recrystallization from the DMSO/chloroform co-solvent system (see Example 3). The XRPD powder pattern of Form IV is shown in **Figure 4** along with some of the 2θ values. Table 4 provides the most prominent 2θ angles, d-spacing and relative intensities for this crystalline material. The XRPD powder pattern for Form IV is distinctly different from that of any other form.

Table 4: 5-azacytidine Form IV-the most prominent 2θ angles, d-spacing and relative intensities (Cu $K\alpha$ radiation)

2θ Angle (°)	d-spacing (Å)	Relative Intensity
5.704	15.408	24.9
11.571	7.642	97.8

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(continued)

	2θ Angle (°)	d-spacing (Å)	Relative Intensity
5	12.563	7.040	22.2
	14.070	6.289	100.0
	15.943	5.555	67.4
	16.993	5.213	51.0
10	18.066	4.906	20.1
	20.377	4.355	44.7
	20.729	4.281	49.0
15	21.484	4.132	36.30
	21.803	4.073	11.2
	22.452	3.957	66.7
	22.709	3.913	64.0
20	23.646	3.760	17.3
	24.068	3.695	19.4
	25.346	3.526	12.0
25	25.346	3.511	12.5
	26.900	3.312	11.0
	27.991	3.185	11.4
	28.527	3.126	25.7
30	28.723	3.106	34.1
	30.124	2.964	14.7
	30.673	2.912	53.6
35	31.059	2.877	15.7
	35.059	2.557	18.1
	38.195	2.354	15.0
40	38.403	2.342	12.6

[0039] Thermal analysis of Form IV is presented in Example 6.

Form V

45 [0040] Form V is a crystalline form of 5-azacytidine. Form V was recovered by fast recrystallization of 5-azacytidine from a DMSO/toluene co-solvent system (see Example 3). The XRPD powder pattern of Form V is shown in Figure 5 along with some of the 2θ values.

Table 5 provides the most prominent 2θ angles, d-spacing and relative intensities for this crystalline material. The XRPD powder pattern for Form V is distinctly different from that of any other form.

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Table 5: 5-azacytidine Form V-the most prominent 2θ angles, d-spacing and relative intensities (Cu Kα radiation)

	2θ Angle (°)	d-spacing (Å)	Relative Intensity
55	11.018	8.024	40.0
	12.351	7.160	29.6
	13.176	6.714	28.3

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(continued)

	2θ Angle (°)	d-spacing (Å)	Relative Intensity
5	13.747	6.436	42.9
	14.548	6.084	18.3
	15.542	5.697	14.2
	16.556	5.350	47.8
10	17.978	4.930	18.1
	18.549	4.780	83.9
	19.202	4.618	25.0
15	19.819	4.476	12.1
	20.329	4.365	28.6
	21.518	4.126	100.0
	21.970	4.042	65.6
20	22.521	3.948	11.5
	23.179	3.834	66.5
	24.018	3.702	13.0
25	24.569	3.620	40.7
	27.224	3.273	50.2
	28.469	3.133	24.2
	29.041	3.072	24.8
30	29.429	3.033	15.0
	30.924	2.889	15.6
	31.133	2.870	22.6
35	37.938	2.370	10.7

[0041] Thermal analysis indicates that Form V is a solvate. See Example 6.

Form VI

40 [0042] The drug product used in the aforementioned NCI investigation was typically prepared by lyophilizing a solution of 5-azacytidine and mannitol (1:1 w/w). The resultant drug product comprised 100 mg of 5-azacytidine and 100 mg mannitol as a lyophilized cake in a vial and was administered by subcutaneous injection as an aqueous suspension ("slurry"). XRPD analysis of retained samples of the drug product used in the NCI investigation revealed the existence of another polymorph, Form VI. The retained drug product samples comprised either Form VI alone, or a Form I/VI mixed phase. Table 6 provides the most prominent 2θ angles, d-spacing and relative intensities for Form VI.

Table 6: 5-azacytidine Form VI - the most prominent 2θ angles, d-spacing and relative intensities (Cu Kα radiation)

	2θ Angle (°)	d-spacing (Å)	Relative Intensity
50	12.533	7.057	10.1
	12.963	6.824	10.2
	13.801	6.411	100.0
55	18.929	4.6843	10.0
	20.920	4.243	34.2

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(continued)

	2θ Angle (°)	d-spacing (Å)	Relative Intensity
5	21.108	4.205	49.4
	21.527	4.125	47.0
	22.623	3.922	10.7
	22.970	3.869	13.8
10	24.054	3.697	77.8
	26.668	3.340	23.0
	27.210	3.275	33.7
15	28.519	3.127	12.9
	29.548	3.021	27.2
	30.458	2.932	50.3
	33.810	2.649	11.6
20	35.079	2.556	12.6
	37.528	2.411	24.7

[0043] Thermal analysis and proton (¹H) NMR spectroscopy of Form VI is presented in Examples 6-7.

Form VII

[0044] Form VII is a crystalline form of 5-azacytidine. Form VII was produced by fast recrystallization from a DM-SO/methanol co-solvent system (see Example 3). Form VII was always isolated by this recrystallization method as a mixed phase with Form I. The XRPD powder pattern of mixed phase Forms I and VII is shown in Figure 7 along with some of the 2θ values and the Form VII distinctive peaks indicated with asterisks. Table 7 provides the most prominent 2θ angles, d-spacing and relative intensities for this mixed phase. Form VII exhibits distinctive peaks at 5.8, 11.5, 12.8, 22.4 and 26.6 ° 2θ in addition to peaks displayed in the Form I XRPD powder pattern. The XRPD pattern for mixed phase Forms I and VII is distinctly different from that of any other form.

Table 7: 5-azacytidine, mixed Forms I and VII the most prominent 2θ angles, d-spacing and relative intensities (Cu Kα radiation)

	2θ Angle (°)	d-spacing (Å)	Relative Intensity
40	5.779	15.281	14.7
	11.537	7.664	8.3
	12.208	7.244	28.0
45	12.759	6.932	21.7
	13.048	6.780	34.4
	14.418	6.138	22.5
	16.489	5.372	21.6
50	18.649	4.754	13.5
	19.101	4.643	34.7
	20.200	4.392	34.4
55	20.769	4.273	10.5
	21.355	4.157	11.7
	22.365	3.972	29.9

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(continued)

2θ Angle (°)	d-spacing (Å)	Relative Intensity
23.049	3.856	100.0
23.884	3.723	23.1
26.628	3.345	13.3
27.145	3.282	52.9
29.296	3.046	26.2
29.582	3.017	11.3
32.078	2.788	12.9

[0045] Thermal analysis of Form VII is presented in Example 6.

Form VIII

[0046] Form VIII is a crystalline form of 5-azacytidine. Form VIII was recovered by recrystallizing 5-azacytidine Form I from a N-methyl-2-pyrrolidone (NMP) single solvent system (see Example 4). The XRPD powder pattern of Form VIII is shown in Figure 8 along with some of the 2θ values. Table 8 provides the most prominent 2θ angles, d-spacing and relative intensities for this material. The XRPD pattern for Form VIII is distinctly different from that of any other form.

Table 8: 5-azacytidine, Form VIII-the most prominent 2θ angles, d-spacing and relative intensities (Cu Kα radiation)

2θ Angle (°)	d-spacing (Å)	Relative Intensity
6.599	13.384	2.9
10.660	8.292	2.2
12.600	7.020	23.4
13.358	6.623	2.6
15.849	5.587	2.0
17.275	5.129	4.2
20.243	4.383	5.8
20.851	4.257	7.8
21.770	4.079	74.4
22.649	3.923	32.1
25.554	3.483	100.0
25.740	3.458	7.8
29.293	3.046	3.8
32.148	2.782	8.8
35.074	2.556	7.4
38.306	2.348	2.5

Amorphous 5-azacytidine

[0047] Amorphous 5-azacytidine may be recovered from equilibrium saturated solutions of 5-azacytidine in propylene glycol, polyethylene glycol and DMSO. See Example 8.

Pharmaceutical Formulations

5 [0048] For the most effective administration of drug substance of the present invention, it is preferred to prepare a pharmaceutical formulation (also known as the "drug product") preferably in unit dose form, comprising one or more of the 5-azacytidine forms of the present invention and one or more pharmaceutically acceptable carrier, diluent, or excipient.

10 [0049] Such pharmaceutical formulation may, without being limited by the teachings set forth herein, include a solid form of the present invention which is blended with at least one pharmaceutically acceptable excipient, diluted by an excipient or enclosed within such a carrier that can be in the form of a capsule, sachet, tablet, buccal, lozenge, paper, or other container. When the excipient serves as a diluent, it may be a solid, semi-solid, or liquid material which acts as a vehicle, carrier, or medium for the 5-azacytidine polymorph(s). Thus, the formulations can be in the form of tablets, pills, powders, elixirs, suspensions, emulsions, solutions, syrups, capsules (such as, for example, soft and hard gelatin capsules), suppositories, sterile injectable solutions, and sterile packaged powders.

15 [0050] Examples of suitable excipients include, but are not limited to, starches, gum arabic, calcium silicate, microcrystalline cellulose, polyvinylpyrrolidone, cellulose, water, syrup, and methyl cellulose. The formulations can additionally include lubricating agents such as, for example, talc, magnesium stearate and mineral oil; wetting agents; emulsifying and suspending agents; preserving agents such as methyl- and propyl- hydroxybenzoates; sweetening agents; or flavoring agents. Polyols, buffers, and inert fillers may also be used. Examples of polyols include, but are not limited to: mannitol, sorbitol, xylitol, sucrose, maltose, glucose, lactose, dextrose, and the like. Suitable buffers encompass, but are not limited to, phosphate, citrate, tartrate, succinate, and the like. Other inert fillers which may be used encompass those which are known in the art and are useful in the manufacture of various dosage forms. If desired, the solid pharmaceutical compositions may include other components such as bulking agents and/or granulating agents, and the like. The compositions of the invention can be formulated so as to provide quick, sustained, controlled, or delayed release of the drug substance after administration to the patient by employing procedures well known in the art.

20 [0051] In certain embodiments of the invention, the 5-azacytidine form(s) may be made into the form of dosage units for oral administration. The 5-azacytidine form(s) may be mixed with a solid, pulverant carrier such as, for example, lactose, saccharose, sorbitol, mannitol, starch, amylopectin, cellulose derivatives or gelatin, as well as with an antifriction agent such as for example, magnesium stearate, calcium stearate, and polyethylene glycol waxes. The mixture is then pressed into tablets or filled into capsules. If coated tablets, capsules, or pulvules are desired, such tablets, capsules, or pulvules may be coated with a concentrated solution of sugar, which may contain gum arabic, gelatin, talc, titanium dioxide, or with a lacquer dissolved in the volatile organic solvent or mixture of solvents. To this coating, various dyes may be added in order to distinguish among tablets with different active compounds or with different amounts of the active compound present.

25 [0052] Soft gelatin capsules may be prepared in which capsules contain a mixture of the 5-azacytidine form(s) and vegetable oil or non-aqueous, water miscible materials such as, for example, polyethylene glycol and the like. Hard gelatin capsules may contain granules or powder of the 5-azacytidine polymorph in combination with a solid, pulverulent carrier, such as, for example, lactose, saccharose, sorbitol, mannitol, potato starch, corn starch, amylopectin, cellulose derivatives, or gelatin.

30 [0053] Tablets for oral use are typically prepared in the following manner, although other techniques may be employed. The solid substances are gently ground or sieved to a desired particle size, and a binding agent is homogenized and suspended in a suitable solvent. The 5-azacytidine form(s) and auxiliary agents are mixed with the binding agent solution. The resulting mixture is moistened to form a uniform suspension. The moistening typically causes the particles to aggregate slightly, and the resulting mass is gently pressed through a stainless steel sieve having a desired size. The layers of the mixture are then dried in controlled drying units for a pre-determined length of time to achieve a desired particle size and consistency. The granules of the dried mixture are gently sieved to remove any powder. To this mixture, disintegrating, anti-friction, and anti-adhesive agents are added. Finally, the mixture is pressed into tablets using a machine with the appropriate punches and dies to obtain the desired tablet size.

35 [0054] In the event that the above formulations are to be used for parenteral administration, such a formulation typically comprises sterile, aqueous and non-aqueous injection solutions comprising one or more 5-azacytidine forms for which preparations are preferably isotonic with the blood of the intended recipient. These preparations may contain anti-oxidants, buffers, bacteriostats, and solute; which render the formulation isotonic with the blood of the intended recipient. Aqueous and non-aqueous suspensions may include suspending agents and thickening agents. The formulations may be present in unit-dose or multi-dose containers, for example, sealed ampules and vials. Extemporaneous injection solutions and suspensions may be prepared from sterile powders, granules, and tablets of the kind previously described.

40 [0055] Liquid preparations for oral administration are prepared in the form of solutions, syrups, or suspensions with the latter two forms containing, for example, 5-azacytidine polymorph(s), sugar, and a mixture of ethanol, water, glycerol, and propylene glycol. If desired, such liquid preparations contain coloring agents, flavoring agents, and saccharin. Thickening agents such as carboxymethylcellulose may also be used.

45 [0056] As such, the pharmaceutical formulations of the present invention are preferably prepared in a unit dosage

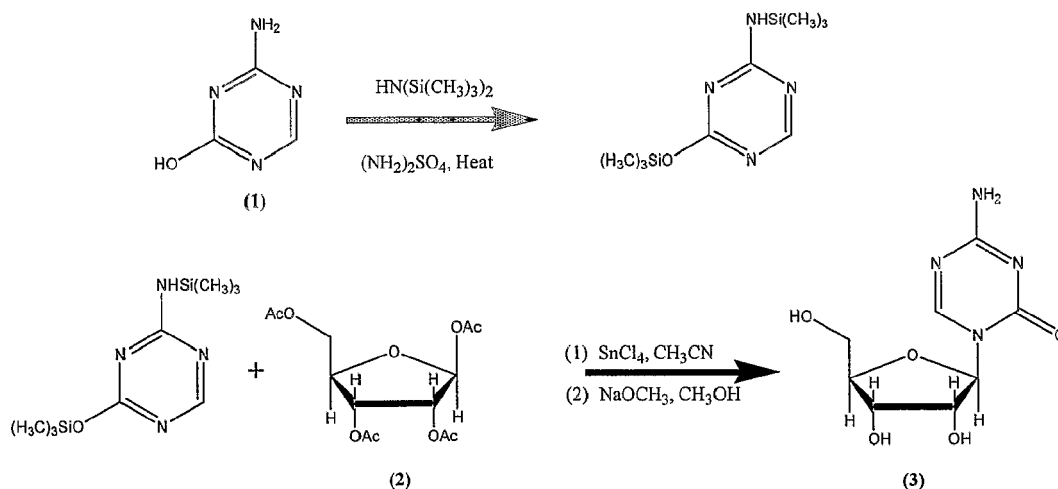
form, each dosage unit containing from about 5 mg to about 200 mg, more usually about 100 mg of the 5-azacytidine form(s). In liquid form, dosage unit contains from about 5 to about 200 mg, more usually about 100 mg of the 5-azacytidine form(s). The term "unit dosage form" refers to physically discrete units suitable as unitary dosages for human subjects/patients or other mammals, each unit containing a predetermined quantity of the 5-azacytidine polymorph calculated to produce the desired therapeutic effect, in association with preferably, at least one pharmaceutically acceptable carrier, diluent, or excipient.

[0057] The following examples are provided for illustrative purposes only, and are not to be construed as limiting the scope of the claims in any way.

Examples

Example 1: Prior Art Procedure for Synthesis of 5-azacytidine Drug Substance

[0058] Using commercially available 5-azacytosine (1) and 1,2,3,5-Tetra-O- β -acetyl-ribofuranose (2) (RTA), 5-azacytidine (3) may be synthesized according to the pathway



below.

[0059] The crude synthesis product is dissolved in DMSO (preheated to about 90°C), and then methanol is added to the DMSO solution. The co-solvent mixture is equilibrated at approximately -20°C to allow 5-azacytidine crystal formation. The product is collected by vacuum filtration and allowed to air dry.

Example 2: Slow Recrystallization from DMSO/toluene

[0060] Dimethyl sulfoxide (DMSO) was used as the primary solvent to solubilize Form I of 5-azacytidine and toluene was used as the co-solvent as follows. Approximately 250 mg of 5-azacytidine was dissolved with approximately 5 mL of DMSO, preheated to approximately 90°C, in separate 100-mL beakers. The solids were allowed to dissolve to a clear solution. Approximately 45 mL of toluene, preheated to approximately 50°C, was added to the solution and the resultant solution was mixed. The solution was covered and allowed to equilibrate at ambient conditions. The product was collected by vacuum filtration as white crystals using a Buchner funnel. The collected product was allowed to air dry.

Example 3: Fast Recrystallization from DMSO/methanol, DMSO/toluene, and DMSO/chloroform

[0061] Approximately 250 mg of 5-azacytidine was dissolved with approximately 5 mL of DMSO as the primary solvent, preheated to approximately 90°C, in separate 100-mL beakers. The solids were allowed to dissolve to a clear solution. Approximately 45 mL of the selected co-solvent (methanol, toluene, or chloroform), preheated to approximately 50°C, was added to the solution and the resultant solution was mixed. The solution was covered and placed in a freezer to equilibrate at approximately -20°C to allow crystal formation. Solutions were removed from the freezer after crystal formation.

[0062] The product from the methanol and toluene solutions was collected by vacuum filtration using a Buchnerfunnel. The resulting white crystalline product was allowed to air dry.

[0063] The chloroform product was too fine to be collected by vacuum filtration. Most of the solvent was carefully

decanted from the chloroform solution and the solvent from the resultant slurry was allowed to evaporate at ambient temperature to dryness. The chloroform solution evaporated to a white product. Note that fast recrystallization using the DMSO/methanol co-solvent system has typically been used to prepare 5-azacytidine drug substance in the prior art (see the last step of the procedure provided in Example 1).

Example 4: Fast Recrystallization N-methyl-2-pyrrolidone (NMP) Single Solvent System

[0064] Approximately 500 mg of 5-azacytidine was dissolved with approximately 5 mL of NMP, preheated to approximately 90 °C, in separate 50-mL beakers. The solids were allowed to dissolve to a clear solution. The solution was covered and placed in a freezer to equilibrate at approximately -20 °C to allow crystal formation. Solutions were removed from the freezer after crystal formation, equilibrated at ambient temperature. The product was collected by vacuum filtration using a Buchner funnel. The collected product was allowed to air dry.

Example 5: X-Ray Powder Diffraction of 5-azacytidine

[0065] X-ray powder diffraction patterns for each sample were obtained on a Scintag XDS 2000 or a Scintag X₂ θ/θ diffractometer operating with copper radiation at 45 kV and 40 mA using a Kevex Psi Peltier-cooled silicon detector or a Thermo ARL Peltier-cooled solid state detector. Source slits of 2 or 4 mm and detector slits of 0.5 or 0.3 mm were used for data collection. Recrystallized material was gently milled using an agate mortar and pestle for approximately one minute. Samples were placed in a stainless steel or silicon sample holder and leveled using a glass microscope slide. Powder diffraction patterns of the samples were obtained from 2 to 42° 2 θ at 1°/minute. Calibration of the X₂ diffractometer is verified annually using a silicon powder standard. Raw data files were converted to ASCII format, transferred to an IBM compatible computer and displayed in Origin® 6.1 for Windows.

[0066] XRPD of a single sample of 5-azacytidine produced according to the method of Example 1 revealed that this sample consisted of Form I of 5-azacytidine.

[0067] NCI retained drug substance sample samples were also analyzed. These samples were all previously synthesized and recrystallized according to the method of Example 1 and were stored at 5°C since production. XRPD revealed some retained samples are comprised of Form I alone, whereas other retained samples contain a mixed phase of Form I and a different polymorph, termed Form II.

[0068] XRPD of NCI retained drug product samples revealed the existence of Form VI in some samples. In those samples, Form VI was present as a mixed phase with Form I.

[0069] XRPD of the recrystallized 5-azacytidine obtained in Example 2 revealed that slow recrystallization from a DMSO/toluene system produced Form IV. XRPD of the recrystallized 5-azacytidine obtained in Example 3 revealed that fast recrystallization from a DMSO/chloroform system produced Form IV, fast recrystallization from a DMSO/toluene system produced Form V, and fast recrystallization from a DMSO/methanol system produced mixed phased Form I/ Form VII. XRPD of the recrystallized 5-azacytidine obtained in Example 4 revealed that the N-methyl-2-pyrrolidone solvent system produced Form VIII.

Example 6: Thermal Analysis of 5-azacytidine

[0070] Differential Scanning Calorimetry (DSC) measurements for each sample were collected using a Perkin Elmer Pyris 1 DSC system equipped with an Intracooler 2P refrigeration unit. The Pyris 1 DSC was purged with nitrogen. Calibration was performed prior to analysis using an Indium standard at a 10 °C minute heating rate. Each sample was gently ground in an agate mortar and pestle. Approximately 1-3 mg of the sample were individually sealed in a Perkin Elmer 30- μ L universal aluminum pan with holes in the lid. Samples were heated from 25 °C to 250 °C or 350 °C at 10 °C/minute.

[0071] Thermogravimetric Analysis (TGA) measurements for each sample were collected using a Perkin Elmer TGA 7 purged with nitrogen at approximately 20 cc/minute. A 100-mg standard weight and nickel metal were used to verify balance and temperature calibrations, respectively. Samples were heated from 25 °C to 250 °C or 300 °C at 10 °C/minute.

[0072] Capillary melting point (MP) measurements were made using an Electrothermal 9300 melting point apparatus. A heating rate of 10 °C/minute was used from set point temperatures described in individual discussions. Visual melting points are reported as an average of triplicate determinations.

[0073] The results are as follows:

Form I

[0074] TGA showed a weight loss of 0.23% between ambient and 150 °C, which indicates that it is anhydrous. DSC exhibited a single event with an onset of 227.0 °C.

[0075] A capillary melting point determination was performed in triplicate on a sample of Form I of 5-azacytidine. The sample was visually observed to decompose without melting at about 215 °C using a 10 °C heating rate and a starting temperature of 200 °C. Thus, the DSC event results from decomposition of 5-azacytidine.

5 *Form III Mixed Phase*

[0076] The TGA for the Form I/II mixed phase showed a weight loss of 1.16% between ambient temperature and 150 °C. The DSC analysis exhibited a single event with an onset at 229.8 °C. The decomposition of the mixed phase was consistent with that observed for 5-azacytidine Form I.

10

Form III

[0077] The TGA showed a weight loss of between 6.56% and 8.44% when the temperature was raised from ambient and 150 °C. The loss is close to the theoretical amount of moisture, 6.9 %, that 5-azacytidine monohydrate would have. The DSC analysis exhibited an endotherm, which is in the range associated with solvent loss, and a higher temperature event. The endotherm exhibited an onset temperature in the range of 86.4-89.2 °C, peak temperatures in the range of 95.8-97.0 °C and ΔH values in the range of 73.1-100.5 J/g. The higher temperature event had onset temperatures in the range 229.1-232.1 °C and was consistent with the decomposition observed for 5-azacytidine Form I.

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[0078] 5-azacytidine Form III was heated at 105 °C for 4 hours in an attempt to dehydrate the material. The material did not change its physical appearance during heating. TGA was used to measure the water content of Form III before and after drying. The initial amount of moisture present in Form III was 6.31 % and was <0.1 % after drying. The XRPD powder pattern for dehydrated Form III matches that of Form I. Thus, Form III dehydrates to Form I.

20

Form IV

[0079] The TGA showed a weight loss of 21.80% between ambient temperature and 150 °C, which does not correspond to the solvent content for any simple solvates. It is not known whether crystalline Form IV is a polymorph or a pseudopolymorph.

25

[0080] The DSC analysis exhibited two endotherms and a higher temperature event. The two endotherms are in the range that is associated with solvent loss. The first endotherm exhibited an onset temperature of 87.6 °C, a peak temperature of 90.1 °C and ΔSH value of 98.3 J/g. The second endotherm exhibited an onset temperature of 136.0 °C, a peak temperature of 139.0 °C and ΔSH value of 81.8 J/g. The higher temperature event had an onset temperature of 230.6 °C and was consistent with the decomposition that was observed for 5-azacytidine Form I.

30

Form V

[0081] TGA showed a weight loss of 21.45% between ambient and 150 °C, which does not correspond to the solvent content for any simple solvate. The DSC analysis exhibited two merged endotherms, a single endotherm and a higher temperature event. The three endotherms are in the range that is associated with solvent loss. The two merged endotherms exhibit onset temperatures of 66.6 and 68.0 °C. The single endotherm exhibited an onset temperature of 88.7 °C, a peak temperature of 121.5 °C and a ΔSH value of 180.3 J/g. The higher temperature event had onset temperature of 230.7 °C and was consistent with the decomposition that was observed for 5-azacytidine Form I.

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Form VI

[0082] TGA showed a weight loss of 1.10% between ambient temperature and 150 °C. The DSC analysis exhibited a small endotherm, an exotherm and a higher temperature event. The small endotherm exhibited an onset temperature of 57.8 °C, a peak temperature of 77.0 °C and a ΔSH value of 55.7 J/g. The exotherm exhibited an onset temperature of 149.3 °C, a peak temperature of 157.1 °C and an ΔSH value of -17.9 J/g. The higher temperature event had an onset temperature of 234.7 °C and was consistent with the decomposition observed for 5-azacytidine Form I.

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Form VII

[0083] TGA showed a weight loss of 2.45% between ambient temperature and 150 °C. The DSC analysis exhibited a minor endotherm and a higher temperature event. The minor endotherm had an onset temperature of 63.3 °C, a peak temperature of 68.3 °C and a ΔSH value of 17.1 J/g. The higher temperature event had an onset temperature of 227.2 °C and is consistent with the decomposition observed for 5-azacytidine Form I.

55

Example 7: Nuclear Magnetic Resonance (NMR) Analysis of Form III and Form VI

[0084] 5-azacytidine is known to be labile in water. Since Form III is found in equilibrium saturated solutions and Form VI is produced by the lyophilization of 5-azacytidine solution, it was of interest to examine the purity of these 5-azacytidine forms using NMR. The proton (¹H) NMR spectra of Form III and Form VI were both consistent with the structure of 5-azacytidine in all essential details.

Example 8: Polymorphic Form Conversion of 5-azacytidine

[0085] Form I of 5-azacytidine was added to various solvents in sufficient quantities to form a slurry, and the slurry allowed to equilibrate for a period of time. The solid material that was present in the slurry was recovered, dried, and analyzed using XRPD (according to the XRPD protocol included in Example 5) with the aim of detecting new polymorphs and pseudopolymorphs during the transition to the dissolved state. Samples equilibrated for 19 hours in saline, 5% dextrose, 5% tween 80, water-saturated octanol, ethanol/water (50/50) and water alone resulted in a distinctly different form of 5-azacytidine, designated Form II (see below). Samples equilibrated for 19 hours in acetone, methyl ethyl ketone, and ethanol resulted in materials identified as Form I. Samples equilibrated for 19 hours in propylene glycol, polyethylene glycol and DMSO resulted in amorphous materials. The results are summarized in Table 9.

Table 9: X-ray Powder Diffraction Analysis Results for Solubility Samples: Form Assignment (Cu K(α) radiation)

Solvent	XRPD Pattern Assignment
Normal Saline	Form III
5% Dextrose	Form III
Acetone	Form I
Propylene glycol	Amorphous
Polyethylene glycol	Amorphous
Methyl ethyl ketone	Form I
5 % Tween 80	Form III
DMSO	Amorphous
Water-saturated Octanol	Form III
Ethyl alcohol	Form I
50/50 EtOH/DI Water	Form III
DI Water	Form III

[0086] The conversion of other forms of 5-azacytidine was also studied. Specifically, a Form I/II mixed phase, Form VI (the lyophilized drug product used in the prior art NCI drug trials), a Form I/VI mixed phase, and a Form I/VII mixed phase were weighed into individual small glass beakers and water was pipeted into each beaker. The sample size and water volume were scaled to maintain an approximate 25 mg/mL ratio. The resultant slurry was allowed to equilibrate for 15 minutes. After equilibration, the sample was filtered and the solid material was dried and analyzed using XRPD. In each case, Form III of 5-azacytidine was observed. The results indicate that all forms of 5-azacytidine convert to Form III during the transition to the dissolved state in water. Thus, when a 5-azacytidine suspension ("slurry") was administered to patients in the aforementioned NCI investigation, the patients received both 5-azacytidine in solution, and Form III of 5-azacytidine.

Claims

1. A pharmaceutical composition comprising a crystalline form of 5-azacytidine for use in a method of treating myelodysplastic syndromes, wherein the crystalline form of 5-azacytidine is **characterized by** peaks observed using X-ray powder diffraction (Cu K α radiation), comprising peaks having the following 2 θ angles:

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2θ Angle (°)
6.566
11.983
13.089
15.138
17.446
20.762
21.049
22.776
24.363
25.743
26.305
28.741
31.393
32.806
33.043
33.536
36.371
39.157
41.643

, and wherein the method comprises the oral administration of the pharmaceutical composition.

2. The pharmaceutical composition for use of claim 1, wherein the crystalline form of 5-azacytidine is **characterized by** an X-ray powder diffraction (Cu Kα radiation) pattern substantially in accordance with Figure 3.
3. The pharmaceutical composition comprising a crystalline form of 5-azacytidine for use of claim 1 or 2, wherein the crystalline form of 5-azacytidine is **characterized by** peaks observed using X-ray powder diffraction (Cu Kα radiation), comprising peaks having the following 2θ angles, d-spacings and relative intensities:

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2θ Angle (°)	d-spacing (Å)	Relative Intensity
6.566	13.450	32.9
11.983	7.380	52.5
13.089	6.758	71.0
15.138	5.848	38.9
17.446	5.079	48.2
20.762	4.275	10.8
21.049	4.147	34.8
22.776	3.901	89.5
24.363	3.651	13.7
25.743	3.458	22.8
26.305	3.385	39.9

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(continued)

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2θ Angle (°)	d-spacing (Å)	Relative Intensity
28.741	3.104	100.0
31.393	2.847	22.5
32.806	2.728	11.8
33.043	2.709	10.1
33.536	2.670	15.1
36.371	2.468	11.0
39.157	2.299	19.3
41.643	2.167	12.1

4. The pharmaceutical composition for use of any of claims 1 to 3, wherein the crystalline form of 5-azacytidine is a monohydrate prepared according to a method comprising:

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contacting a crystalline form of 5-azacytidine with water; and
isolating the crystalline form of 5-azacytidine.

5. The pharmaceutical composition for use of claim 4, wherein the crystalline form of 5-azacytidine used in the preparation of the crystalline form 5-azacytidine according to claims 1 to 3 is Form I, Form II, Form IV, Form V, Form VI, Form VII, or Form VIII of 5-azacytidine; or a mixture of two or more thereof.

6. A pharmaceutical composition comprising a mixed phase of forms of 5-azacytidine for use in a method of treating myelodysplastic syndromes, wherein the mixed phase of forms of 5-azacytidine is **characterized by** peaks observed using X-ray powder diffraction (Cu Kα radiation), having the following 2θ angles:

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2θ Angle (°)
12.244
13.082
13.458
14.452
16.521
17.648
18.677
19.093
20.231
21.353
22.309
23.070
23.909
26.641
26.813
27.158
29.309
29.609

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(continued)

2θ Angle (°)
30.384
32.074

, and wherein the method comprises the oral administration of the pharmaceutical composition.

7. The pharmaceutical composition for use of claim 6, wherein the mixed phase of forms of 5-azacytidine is **characterized by** an X-ray powder diffraction pattern substantially in accordance with Figure 2.
8. The pharmaceutical composition for use of claims 6 or 7, wherein the mixed phase of forms of 5-azacytidine is **characterized by** peaks observed using X-ray powder diffraction (Cu K α radiation), comprising peaks having the following 2θ angles, d-spacings and relative intensities:

2θ Angle (°)	d-spacing (Å)	Relative Intensity
12.244	7.223	34.8
13.082	6.762	37.0
13.458	6.574	29.2
14.452	6.124	25.4
16.521	5.361	19.0
17.648	5.022	12.1
18.677	4.747	12.7
19.093	4.645	41.3
20.231	4.386	42.1
21.353	4.158	15.5
22.309	3.982	35.1
23.070	3.852	100.0
23.909	3.719	18.9
26.641	3.343	18.2
26.813	3.322	12.6
27.158	3.281	46.0
29.309	3.045	27.3
29.609	3.015	12.7
30.384	2.939	10.5
32.074	2.788	12.0

9. A pharmaceutical composition comprising a crystalline form of 5-azacytidine for use in a method of treating myelodysplastic syndromes, wherein the crystalline form is **characterized by** peaks observed using X-ray powder diffraction (Cu K α radiation), comprising peaks having the following 2θ angles:

2θ Angle (°)
12.533
12.963

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(continued)

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2θ Angle (°)
13.801
18.929
20.920
21.108
21.527
22.623
22.970
24.054
26.668
27.210
28.519
29.548
30.458
33.810
35.079
37.528

, and wherein the method comprises the oral administration of the pharmaceutical composition.

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10. The pharmaceutical composition for use of claim 9, wherein the crystalline form of 5-azacytidine is **characterized by** an X-ray powder diffraction (Cu Kα radiation) pattern substantially in accordance with Figure 6.
11. The pharmaceutical composition comprising a crystalline form of 5-azacytidine for use of claim 9 or 10, wherein the crystalline form of 5-azacytidine is **characterized by** peaks observed using X-ray powder diffraction (Cu Kα radiation), comprising peaks having the following 2θ angles, d-spacings and relative intensities:

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2θ Angle (°)	d-spacing (Å)	Relative Intensity
12.533	7.057	10.1
12.963	6.824	10.2
13.801	6.411	100.0
18.929	4.6843	10.0
20.920	4.243	34.2
21.108	4.205	49.4
21.527	4.125	47.0
22.623	3.922	10.7
22.970	3.869	13.8
24.054	3.697	77.8
26.668	3.340	23.0
27.210	3.275	33.7
28.519	3.127	12.9

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(continued)

2θ Angle (°)	d-spacing (Å)	Relative Intensity
29.548	3.021	27.2
30.458	2.932	50.3
33.810	2.649	11.6
35.079	2.556	12.6
37.528	2.411	24.7

12. The pharmaceutical composition for use of any one of claims 9 to 11, wherein the crystalline form of 5-azacytidine is prepared according to a method comprising:

lyophilizing a solution of 5-azacytidine and mannitol; and
isolating the lyophilized solid.

13. The pharmaceutical composition for use of claim 12, wherein the solution of 5-azacytidine and mannitol comprises 5-azacytidine and mannitol in a ratio of about 1:1 by weight.

14. A pharmaceutical composition comprising an amorphous solid of 5-azacytidine for use in a method of treating myelodysplastic syndromes, wherein the method comprises the oral administration of the pharmaceutical composition.

15. The pharmaceutical composition for use of claim 14, wherein the amorphous solid of 5-azacytidine is prepared by a method comprising:

adding a crystalline form of 5-azacytidine, being **characterized by** peaks observed using X-ray powder diffraction (Cu K α radiation) having the following 2θ angles:

2θ Angle (°)
12.182
13.024
14.399
16.470
18.627
19.049
20.182
21.329
23.033
23.872
26.863
27.135
29.277
29.591
30.369
32.072

, to a solvent selected from the group consisting of propylene glycol, polyethylene glycol, and DMSO;

allowing equilibration to occur; and recovering 5-azacytidine therefrom.

5 16. The pharmaceutical composition for use of any of the preceding claims, wherein the pharmaceutical composition further comprises a pharmaceutically acceptable excipient, diluent, or carrier; or wherein the pharmaceutical composition is made in an unit dosage form.

10 17. The pharmaceutical composition for use of claim 16, wherein each dosage unit contains from 5 mg to 200 mg of the 5-azacytidine form, preferably 100 mg of the 5-azacytidine form.

Patentansprüche

15 1. Pharmazeutische Zusammensetzung umfassend eine kristalline Form von 5-Azacytidin für die Verwendung in einem Verfahren zur Behandlung von myelodysplastischen Syndromen, wobei die kristalline Form von 5-Azacytidin durch Peaks charakterisiert ist, die unter Verwendung von Röntgenpulverdiffraktometrie (Cu K α Strahlung) beobachtet werden, umfassend Peaks mit den folgenden 2 θ Winkeln:

2 θ Winkel (°)
6.566
11.983
13.089
15.138
17.446
20.762
21.049
22.776
24.363
25.743
26.305
28.741
31.393
32.806
33.043
33.536
36.371
39.157
41.643

20 und wobei das Verfahren die orale Verabreichung der pharmazeutischen Zusammensetzung umfasst.

25 2. Pharmazeutische Zusammensetzung für die Verwendung nach Anspruch 1, wobei die kristalline Form von 5-Azacytidin durch ein Röntgenpulverdiffraktometriemuster (Cu K α Strahlung), das im Wesentlichen mit Figur 3 übereinstimmt, charakterisiert ist.

30 3. Pharmazeutische Zusammensetzung umfassend eine kristalline Form von 5-Azacytidin für die Verwendung nach Anspruch 1 oder 2, wobei die kristalline Form von 5-Azacytidin durch Peaks charakterisiert ist, die unter Verwendung von Röntgenpulverdiffraktometrie (Cu K α Strahlung) beobachtet werden, umfassend Peaks mit den folgenden 2 θ

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Winkeln, d-Abständen und relativen Intensitäten:

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20
25
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2θ Winkel (°)	d-Abstand (Å)	Relative Intensität
6.566	13.450	32.9
11.983	7.380	52.5
13.089	6.758	71.0
15.138	5.848	38.9
17.446	5.079	48.2
20.762	4.275	10.8
21.049	4.147	34.8
22.776	3.901	89.5
24.363	3.651	13.7
25.743	3.458	22.8
26.305	3.385	39.9
28.741	3.104	100.0
31.393	2.847	22.5
32.806	2.728	11.8
33.043	2.709	10.1
33.536	2.670	15.1
36.371	2.468	11.0
39.157	2.299	19.3
41.643	2.167	12.1

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4. Pharmazeutische Zusammensetzung für die Verwendung nach einem der Ansprüche 1 bis 3, wobei die kristalline Form von 5-Azacytidin ein Monohydrat ist, das gemäß einem Verfahren hergestellt wird umfassend:

In-Kontakt-bringen einer kristallinen Form von 5-Azacytidin mit Wasser; und
Isolieren der kristallinen Form von 5-Azacytidin.

5. Pharmazeutische Zusammensetzung für die Verwendung nach Anspruch 4, wobei die kristalline Form von 5-Azacytidin, die bei der Herstellung der kristallinen Form von 5-Azacytidin gemäß den Ansprüchen 1 bis 3 verwendet wird, Form I, Form II, Form IV, Form V, Form VI, Form VII oder Form VIII von 5-Azacytidin ist; oder eine Mischung von zwei oder mehreren davon.

6. Pharmazeutische Zusammensetzung umfassend eine Mischphase von Formen von 5-Azacytidin für die Verwendung in einem Verfahren zur Behandlung von myelodysplastischen Syndromen, wobei die Mischphase von Formen von 5-Azacytidin durch Peaks, die unter Verwendung von Röntgenpulverdiffraktometrie (Cu K α Strahlung) beobachtet werden, mit den folgenden 2 θ Winkeln charakterisiert ist:

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2θ Winkel (°)
12.244
13.082
13.458
14.452
16.521

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(fortgesetzt)

5
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2 θ Winkel (°)
17.648
18.677
19.093
20.231
21.353
22.309
23.070
23.909
26.641
26.813
27.158
29.309
29.609
30.384
32.074

und wobei das Verfahren die orale Verabreichung der pharmazeutischen Zusammensetzung umfasst.

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7. Pharmazeutische Zusammensetzung für die Verwendung nach Anspruch 6, wobei die Mischphase von Formen von 5-Azacytidin durch ein Röntgenpulverdiffraktometrimuster (Cu K α Strahlung), das im Wesentlichen mit Figur 2 übereinstimmt, charakterisiert ist.
8. Pharmazeutische Zusammensetzung für die Verwendung nach den Ansprüchen 6 oder 7, wobei die Mischphase von Formen von 5-Azacytidin durch Peaks charakterisiert ist, die unter Verwendung von Röntgenpulverdiffraktometrie (Cu K α Strahlung) beobachtet werden, umfassend Peaks mit den folgenden 2 θ Winkeln, d-Abständen und relative Intensitäten:

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2 θ Winkel (°)	d-Abstand (Å)	Relative Intensität
12.244	7.223	34.8
13.082	6.762	37.0
13.458	6.574	29.2
14.452	6.124	25.4
16.521	5.361	19.0
17.648	5.022	12.1
18.677	4.747	12.7
19.093	4.645	41.3
20.231	4.386	42.1
21.353	4.158	15.5
22.309	3.982	35.1
23.070	3.852	100.0
23.909	3.719	18.9

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(fortgesetzt)

2 θ Winkel (°)	d-Abstand (Å)	Relative Intensität
26.641	3.343	18.2
26.813	3.322	12.6
27.158	3.281	46.0
29.309	3.045	27.3
29.609	3.015	12.7
30.384	2.939	10.5
32.074	2.788	12.0

9. Pharmazeutische Zusammensetzung umfassend eine kristalline Form von 5-Azacytidin für die Verwendung in einem Verfahren zur Behandlung von myelodysplastischen Syndromen, wobei die kristalline Form durch Peaks charakterisiert ist, die unter Verwendung von Röntgenpulverdiffraktometrie (Cu K α Strahlung) beobachtet werden, umfassend Peaks mit den folgenden 2 θ Winkeln:

2 θ Winkel (°)
12.533
12.963
13.801
18.929
20.920
21.108
21.527
22.623
22.970
24.054
26.668
27.210
28.519
29.548
30.458
33.810
35.079
37.528

und wobei das Verfahren die orale Verabreichung der pharmazeutischen Zusammensetzung umfasst.

10. Pharmazeutische Zusammensetzung für die Verwendung nach Anspruch 9, wobei die kristalline Form von 5-Azacytidin durch ein Röntgenpulverdiffraktometriemuster (Cu K α Strahlung), das im Wesentlichen mit Figur 6 übereinstimmt, charakterisiert ist,
11. Pharmazeutische Zusammensetzung umfassend eine kristalline Form von 5-Azacytidin für die Verwendung nach Anspruch 9 oder 10, wobei die kristalline Form von 5-Azacytidin durch Peaks charakterisiert ist, die unter Verwendung von Röntgenpulverdiffraktometrie (Cu K α Strahlung) beobachtet werden, umfassend Peaks mit den folgenden 2 θ

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Winkeln, d-Abständen und relativen Intensitäten:

5
10
15
20
25
30

2θ Winkel (°)	d-Abstand (Å)	Relative Intensität
12.533	7.057	10.1
12.963	6.824	10.2
13.801	6.411	100.0
18.929	4.6843	10.0
20.920	4.243	34.2
21.108	4.205	49.4
21.527	4.125	47.0
22.623	3.922	10.7
22.970	3.869	13.8
24.054	3.697	77.8
26.668	3.340	23.0
27.210	3.275	33.7
28.519	3.127	12.9
29.548	3.021	27.2
30.458	2.932	50.3
33.810	2.649	11.6
35.079	2.556	12.6
37.528	2.411	24.7

12. Pharmazeutische Zusammensetzung für die Verwendung nach einem der Ansprüche 9 bis 11, wobei die kristalline Form von 5-Azacytidin gemäß einem Verfahren hergestellt wird umfassend:

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Lyophilisieren einer Lösung von 5-Azacytidin und Mannitol; und Isolieren des lyophilisierten Feststoffs.

13. Pharmazeutische Zusammensetzung für die Verwendung nach Anspruch 12, wobei die Lösung von 5-Azacytidin und Mannitol 5-Azacytidin und Mannitol in einem Gewichtsverhältnis von ungefähr 1:1 umfasst.

14. Pharmazeutische Zusammensetzung umfassend einen amorphen Feststoff von 5-Azacytidin für die Verwendung in einem Verfahren zur Behandlung von myelodysplastischen Syndromen, wobei das Verfahren die orale Verabreichung der pharmazeutischen Zusammensetzung umfasst.

15. Pharmazeutische Zusammensetzung für die Verwendung nach Anspruch 14, wobei der amorphe Feststoff von 5-Azacytidin durch ein Verfahren hergestellt wird umfassend:

50 Hinzufügen einer kristallinen Form von 5-Azacytidin, die durch Peaks, die unter Verwendung von Röntgenpulverdiffraktometrie (Cu K α Strahlung) beobachtet werden, mit den folgenden 2 θ Winkeln charakterisiert ist:

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2θ Winkel (°)
12.182
13.024
14.399
16.470

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(fortgesetzt)

2 θ Winkel (°)
18.627
19.049
20.182
21.329
23.033
23.872
26.863
27.135
29.277
29.591
30.369
32.072

zu einem Lösungsmittel ausgewählt aus der Gruppe bestehend aus Propylenglycol, Polyethylenglycol und DMSO;

Ermöglichen der Gleichgewichtseinstellung; und Wiedergewinnen von 5-Azacytidin davon.

16. Pharmazeutische Zusammensetzung für die Verwendung nach einem der vorangegangenen Ansprüche, wobei die pharmazeutische Zusammensetzung weiter einen pharmazeutisch annehmbaren Exzipienten, Verdünnungsmittel oder Träger umfasst; oder wobei die pharmazeutische Zusammensetzung als eine Einzeldosierungsform hergestellt ist.

17. Pharmazeutische Zusammensetzung für die Verwendung nach Anspruch 16, wobei jede Einzeldosierung 5 mg bis 200 mg der Form von 5-Azacytidin, bevorzugt 100 mg der Form von 5-Azacytidin enthält.

Revendications

1. Composition pharmaceutique comprenant une forme cristalline de la 5-azacytidine pour une utilisation dans une méthode de traitement des syndromes myélodysplasiques, dans laquelle la forme cristalline de la 5-azacytidine est **caractérisée par** des pics observés au moyen d'une diffraction des rayons X sur poudre (rayonnement $K\alpha$ du Cu), comprenant les angles 2θ suivants :

Angle 2 θ (°)
6,566
11,983
13,089
15,138
17,446
20,762
21,049
22,776
24,363

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(suite)

Angle 2 θ (°)
25,743
26,305
28,741
31,393
32,806
33,043
33,536
36,371
39,157
41,643

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, et dans laquelle la méthode comprend l'administration orale de la composition pharmaceutique.

2. Composition pharmaceutique pour une utilisation selon la revendication 1, dans laquelle la forme cristalline de la 5-azacytidine est **caractérisée par** un diagramme de diffraction des rayons X sur poudre (rayonnement K α du Cu) sensiblement selon la figure 3.

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3. Composition pharmaceutique comprenant une forme cristalline de la 5-azacytidine pour une utilisation selon la revendication 1 ou 2, dans laquelle la forme cristalline de la 5-azacytidine est **caractérisée par** des pics observés au moyen d'une diffraction des rayons X sur poudre (rayonnement K α du Cu) comprenant des pics ayant les angles 2 θ , les espacements d et les intensités relatives qui suivent :

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Angle 2 θ (°)	Espacements d (Å)	Intensité relative
6,566	13,450	32,9
11,983	7,380	52,5
13,089	6,758	71,0
15,138	5,848	38,9
17,446	5,079	48,2
20,762	4,275	10,8
21,049	4,147	34,8
22,776	3,901	89,5
24,363	3,651	13,7
25,743	3,458	22,8
26,305	3,385	39,9
28,741	3,104	100,0
31,393	2,847	22,5
32,806	2,728	11,8
33,043	2,709	10,1
33,536	2,670	15,1
36,371	2,468	11,0
39,157	2,299	19,3

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(suite)

Angle 2 θ (°)	Espacements d (Å)	Intensité relative
41,643	2,167	12,1

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4. Composition pharmaceutique pour une utilisation selon l'une quelconque des revendications 1 à 3, dans laquelle la forme cristalline de la 5-azacytidine est un monohydrate préparé selon un procédé comprenant :

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la mise en contact d'une forme cristalline de la 5-azacytidine avec de l'eau ; et
l'isolement de la forme cristalline de 5-azacytidine.

5. Composition pharmaceutique pour une utilisation selon la revendication 4, dans laquelle la forme cristalline de la 5-azacytidine utilisée dans la préparation de la forme cristalline de 5-azacytidine selon les revendications 1 à 3 est la forme I, la forme II, la forme IV, la forme V, la forme VI, la forme VII ou la forme VIII de la 5-azacytidine ; ou un mélange de deux ou plus de celles-ci.

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6. Composition pharmaceutique comprenant une phase mixte de formes de la 5-azacytidine pour une utilisation dans une méthode de traitement des syndromes myélodysplasiques, dans laquelle la phase mixte de formes de la 5-azacytidine est **caractérisée par** des pics observés au moyen d'une diffraction des rayons X sur poudre (rayonnement K α du Cu), ayant les angles 2 θ suivants :

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Angle 2 θ (°)
12,244
13,082
13,458
14,452
16,521
17,648
18,677
19,093
20,231
21,353
22,309
23,070
23,909
26,641
26,813
27,158
29,309
29,609
30,384
32,074

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, et dans laquelle la méthode comprend l'administration orale de la composition pharmaceutique.

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7. Composition pharmaceutique pour une utilisation selon la revendication 6, dans laquelle la phase mixte de formes de la 5-azacytidine est **caractérisée par** un diagramme de diffraction des rayons X sur poudre sensiblement selon

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la figure 2.

8. Composition pharmaceutique pour une utilisation selon les revendications 6 ou 7, dans laquelle la phase mixte de formes la de 5-azacytidine est **caractérisée par** des pics observés au moyen d'une diffraction des rayons X sur poudre (rayonnement $K\alpha$ du Cu), comprenant des pics ayant les angles 2θ , les espacements d et les intensités relatives qui suivent :

Angle 2θ (°)	Espacements d (Å)	Intensité relative
12,244	7,223	34,8
13,082	6,762	37,0
13,458	6,574	29,2
14,452	6,124	25,4
16,521	5,361	19,0
17,648	5,022	12,1
18,677	4,747	12,7
19,093	4,645	41,3
20,231	4,386	42,1
21,353	4,158	15,5
22,309	3,982	35,1
23,070	3,852	100,0
23,909	3,719	18,9
26,641	3,343	18,2
26,813	3,322	12,6
27,158	3,281	46,0
29,309	3,045	27,3
29,609	3,015	12,7
30,384	2,939	10,5
32,074	2,788	12,0

9. Composition pharmaceutique comprenant une forme cristalline de la 5-azacytidine pour une utilisation dans une méthode de traitement des syndromes myélodysplasiques, dans laquelle la forme cristalline est **caractérisée par** des pics observés au moyen d'une diffraction des rayons X sur poudre (rayonnement $K\alpha$ du Cu), comprenant des pics ayant les angles 2θ suivants :

Angle 2θ (°)
12,533
12,963
13,801
18,929
20,920
21,108
21,527
22,623

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(suite)

Angle 2 θ (°)
22,970
24,054
26,668
27,210
28,519
29,548
30,458
33,810
35,079
37,528

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, et dans laquelle la méthode comprend l'administration orale de la composition pharmaceutique.

10. Composition pharmaceutique pour une utilisation selon la revendication 9, dans laquelle la forme cristalline de la 5-azacytidine est **caractérisée par** un diagramme de diffraction des rayons X sur poudre (rayonnement K α du Cu) sensiblement selon la figure 6.

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11. Composition pharmaceutique comprenant une forme cristalline de la 5-azacytidine pour une utilisation selon la revendication 9 ou 10, dans laquelle la forme cristalline de la 5-azacytidine est **caractérisée par** des pics observés au moyen d'une diffraction des rayons X sur poudre (rayonnement K α du Cu), comprenant des pics ayant les angles 2 θ , les espacements d et les intensités relatives qui suivent :

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Angle 2 θ (°)	Espacements d (Å)	Intensité relative
12,533	7,057	10,1
12,963	6,824	10,2
13,801	6,411	100,0
18,929	4,6843	10,0
20,920	4,243	34,2
21,108	4,205	49,4
21,527	4,125	47,0
22,623	3,922	10,7
22,970	3,869	13,8
24,054	3,697	77,8
26,668	3,340	23,0
27,210	3,275	33,7
28,519	3,127	12,9
29,548	3,021	27,2
30,458	2,932	50,3
33,810	2,649	11,6
35,079	2,556	12,6
37,528	2,411	24,7

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12. Composition pharmaceutique pour une utilisation selon l'une quelconque des revendications 9 à 11, dans laquelle la forme cristalline de la 5-azacytidine est préparée selon un procédé comprenant :

la lyophilisation d'une solution de 5-azacytidine et de mannitol ; et
l'isolement du solide lyophilisé.

13. Composition pharmaceutique pour une utilisation selon la revendication 12, dans laquelle la solution de 5-azacytidine et de mannitol comprend de la 5-azacytidine et du mannitol dans un rapport d'environ 1:1 en poids.

14. Composition pharmaceutique comprenant un solide amorphe de 5-azacytidine pour une utilisation dans une méthode de traitement des syndromes myélodysplasiques, dans laquelle la méthode comprend l'administration orale de la composition pharmaceutique.

15. Composition pharmaceutique pour une utilisation selon la revendication 14, dans laquelle le solide amorphe de 5-azacytidine est préparé par un procédé comprenant :

l'ajout d'une forme cristalline de la 5-azacytidine, étant **caractérisée par** des pics observés au moyen d'une diffraction des rayons X sur poudre (rayonnement $K\alpha$ du Cu) ayant les angles 2θ suivants :

Angle 2θ (°)
12,182
13,024
14,399
16,470
18,627
19,049
20,182
21,329
23,033
23,872
26,863
27,135
29,277
29,591
30,369
32,072

, à un solvant choisi dans le groupe constitué par le propylène glycol, le polyéthylène glycol et le DMSO ;
permettre à l'équilibration de se produire ; et
la récupération de la 5-azacytidine à partir de celui-ci.

16. Composition pharmaceutique pour une utilisation selon l'une quelconque des revendications précédentes, dans laquelle la composition pharmaceutique comprend en outre un excipient, un diluant ou un support pharmaceutiquement acceptable ; ou dans laquelle la composition pharmaceutique est réalisée sous une forme posologique unitaire.

17. Composition pharmaceutique pour une utilisation selon la revendication 16, dans laquelle chaque unité posologique contient de 5 mg à 200 mg de la forme de 5-azacytidine, de préférence 100 mg de la forme de 5-azacytidine.

X-ray Powder Diffraction Pattern of Azacitidine, Form I, Labeled with the more Prominent 2 θ Angles (Cu K α Radiation)

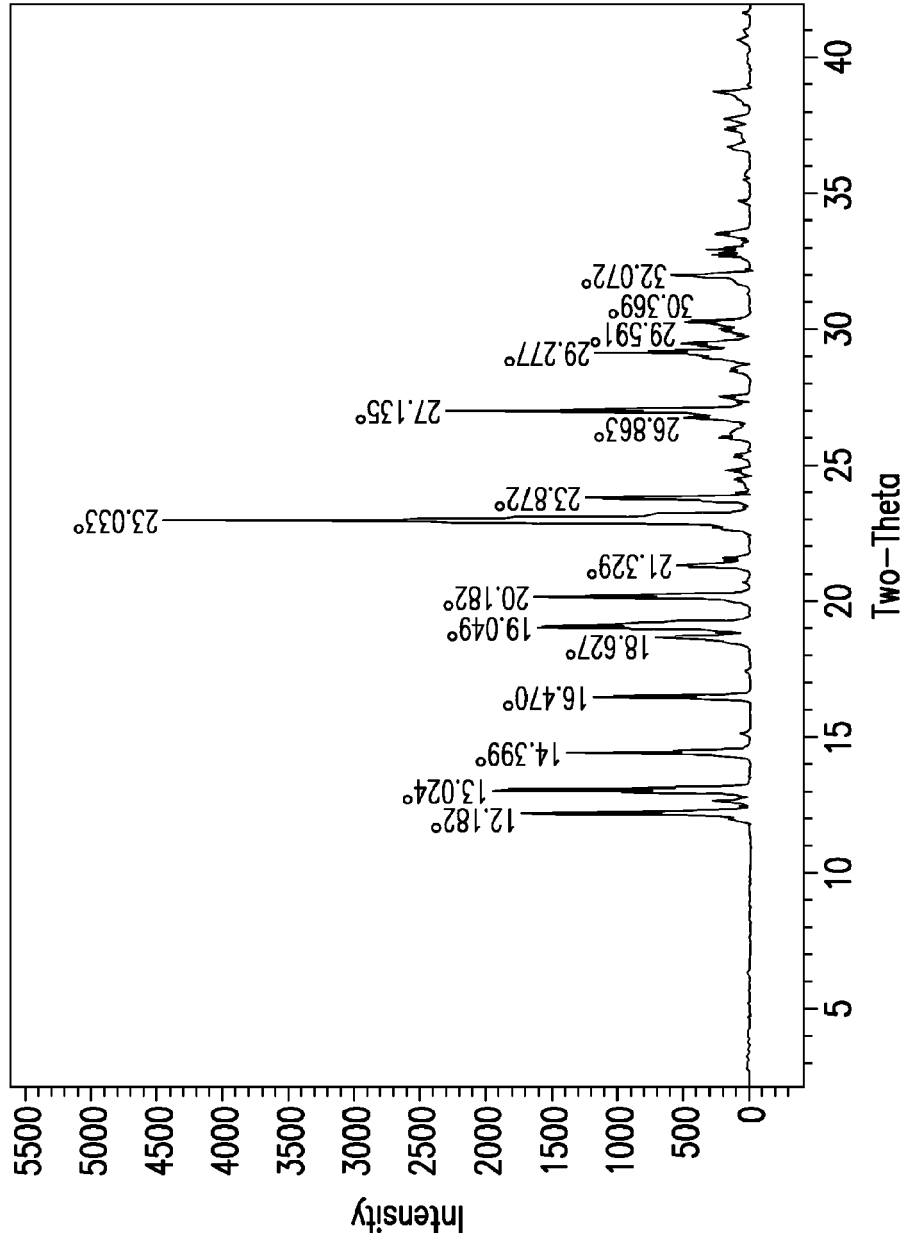


FIG.1

X-ray Powder Diffraction Pattern of Azacitidine, Mixed Phase Forms I and II, Labeled with the more Prominent 2 θ Angles (Cu K α Radiation)

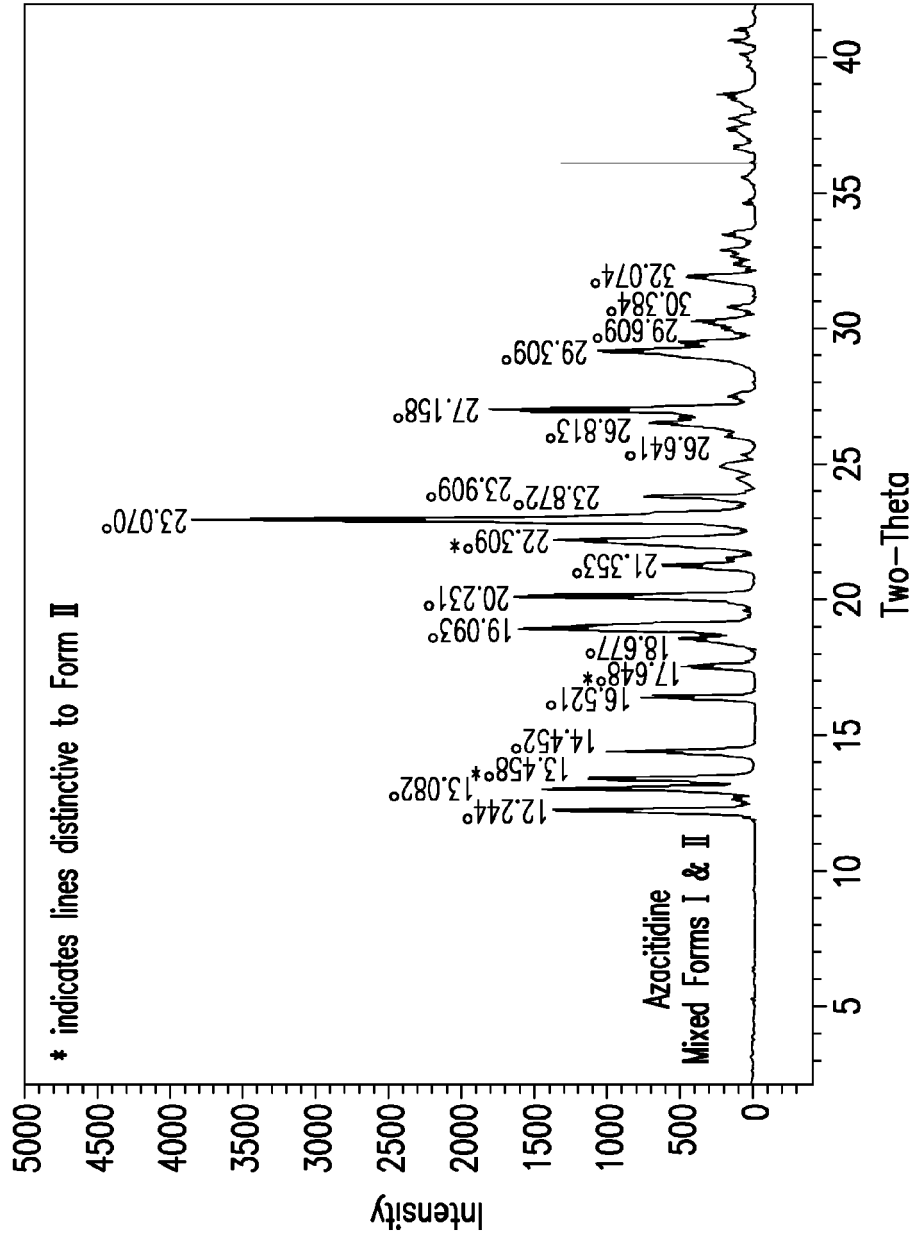


FIG.2

X-ray Powder Diffraction Pattern of Azacitidine, Form III, Labeled with the more Prominent 2 θ Angles (Cu K α Radiation)

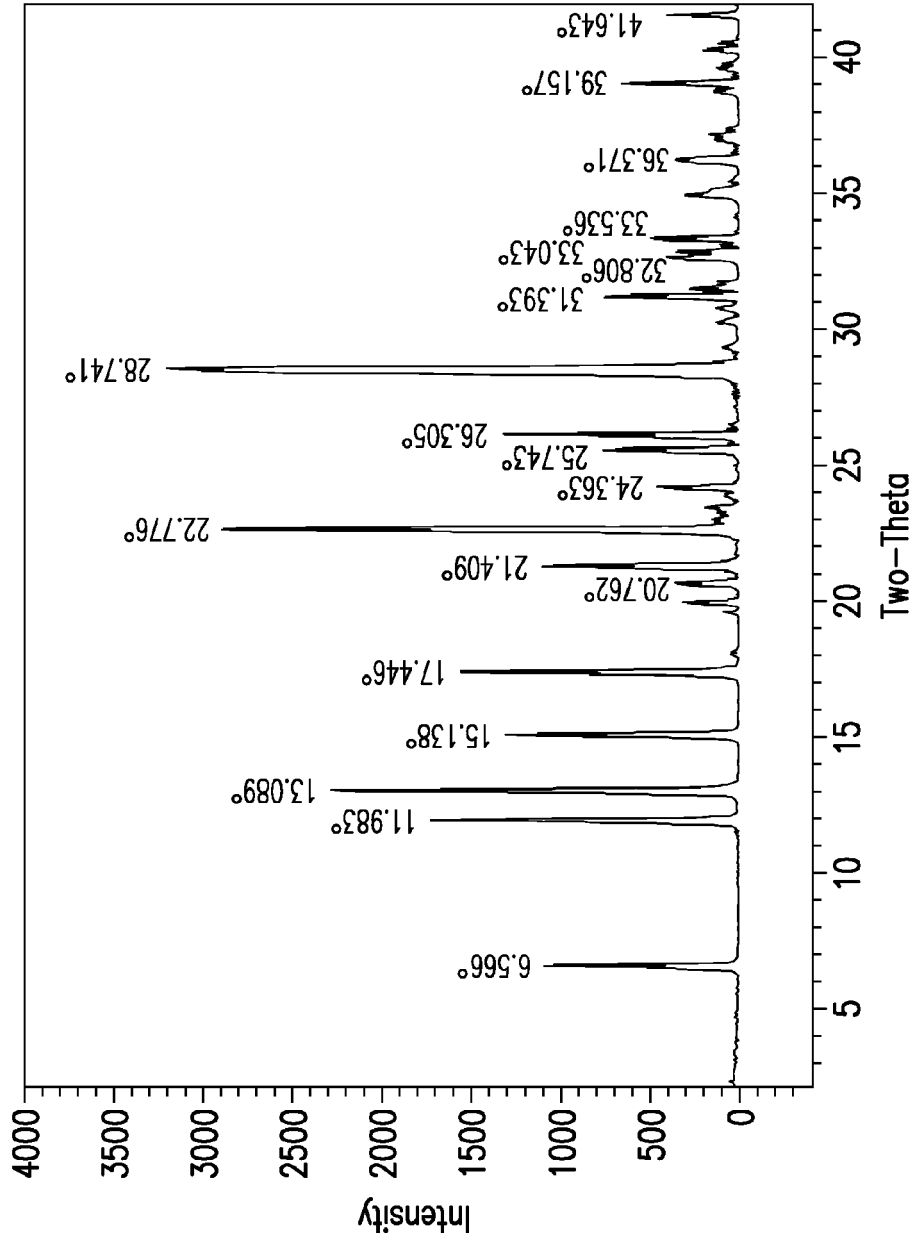


FIG.3

X-ray Powder Diffraction Pattern of Azacitidine, Form IV, Labeled with the more Prominent 2 θ Angles (Cu K α Radiation)

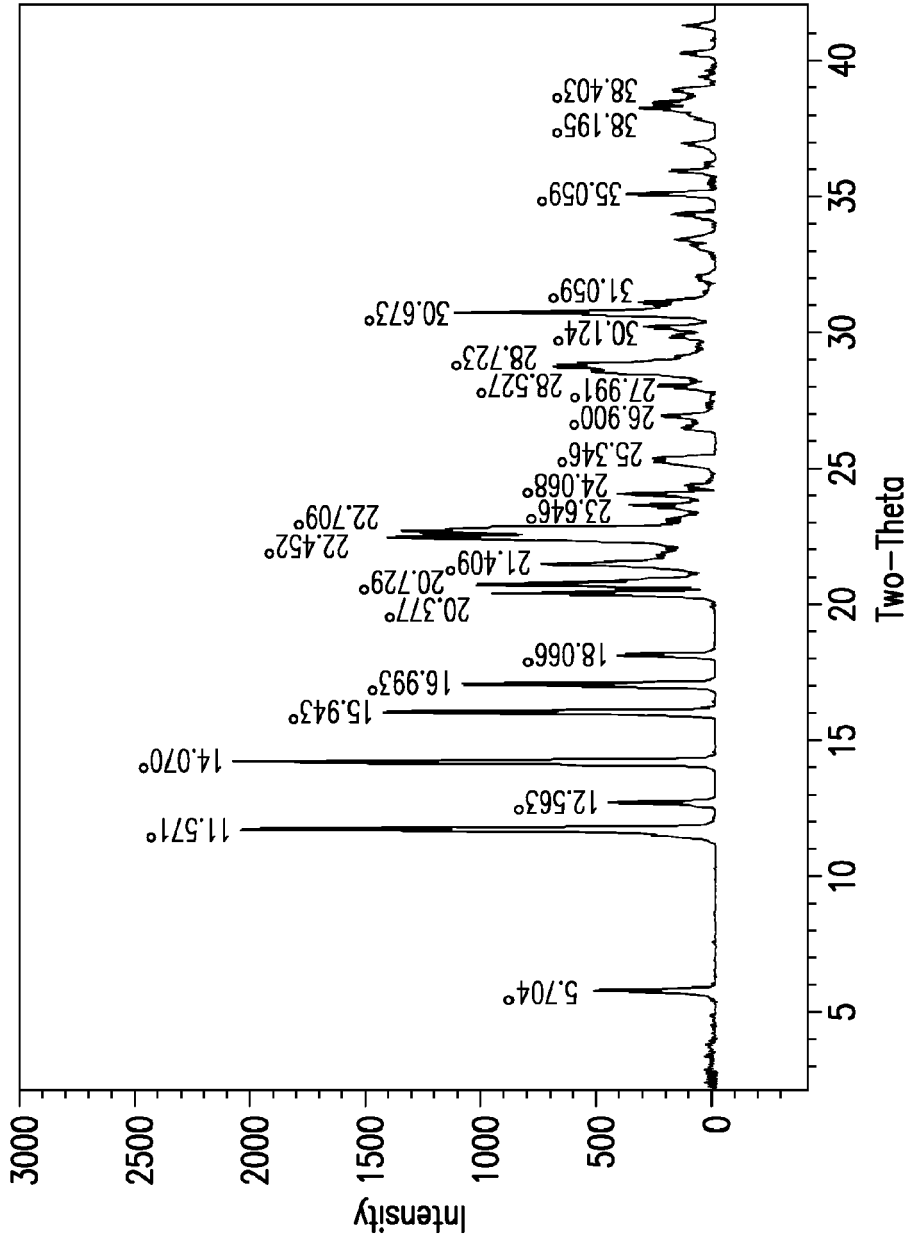


FIG.4

X-ray Powder Diffraction Pattern of Azacitidine, Form V, Labeled with the more Prominent 2 θ Angles (Cu K α Radiation)

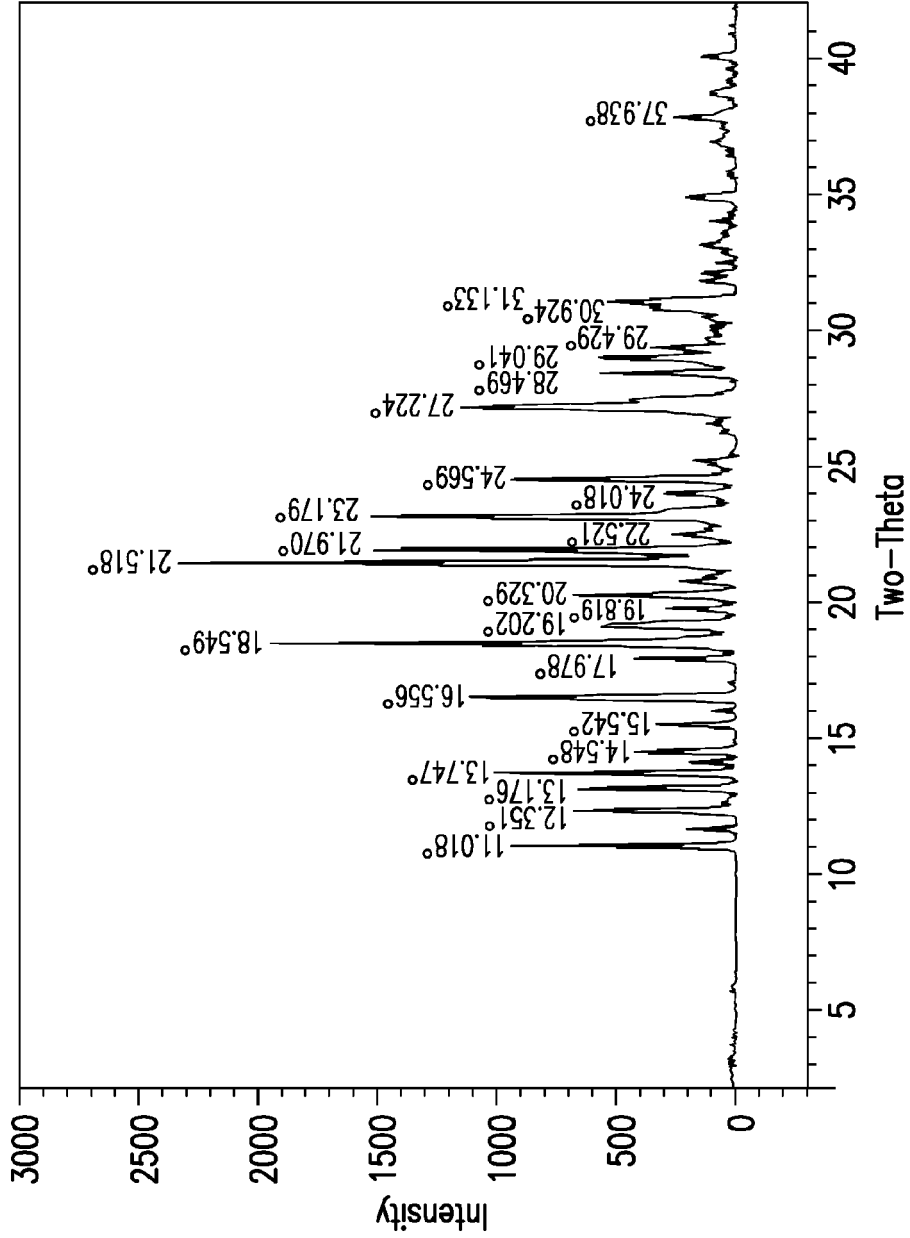


FIG.5

X-ray Powder Diffraction Pattern of Azacitidine, Form VI, Labeled with the more Prominent 2 θ Angles (Cu K α Radiation)

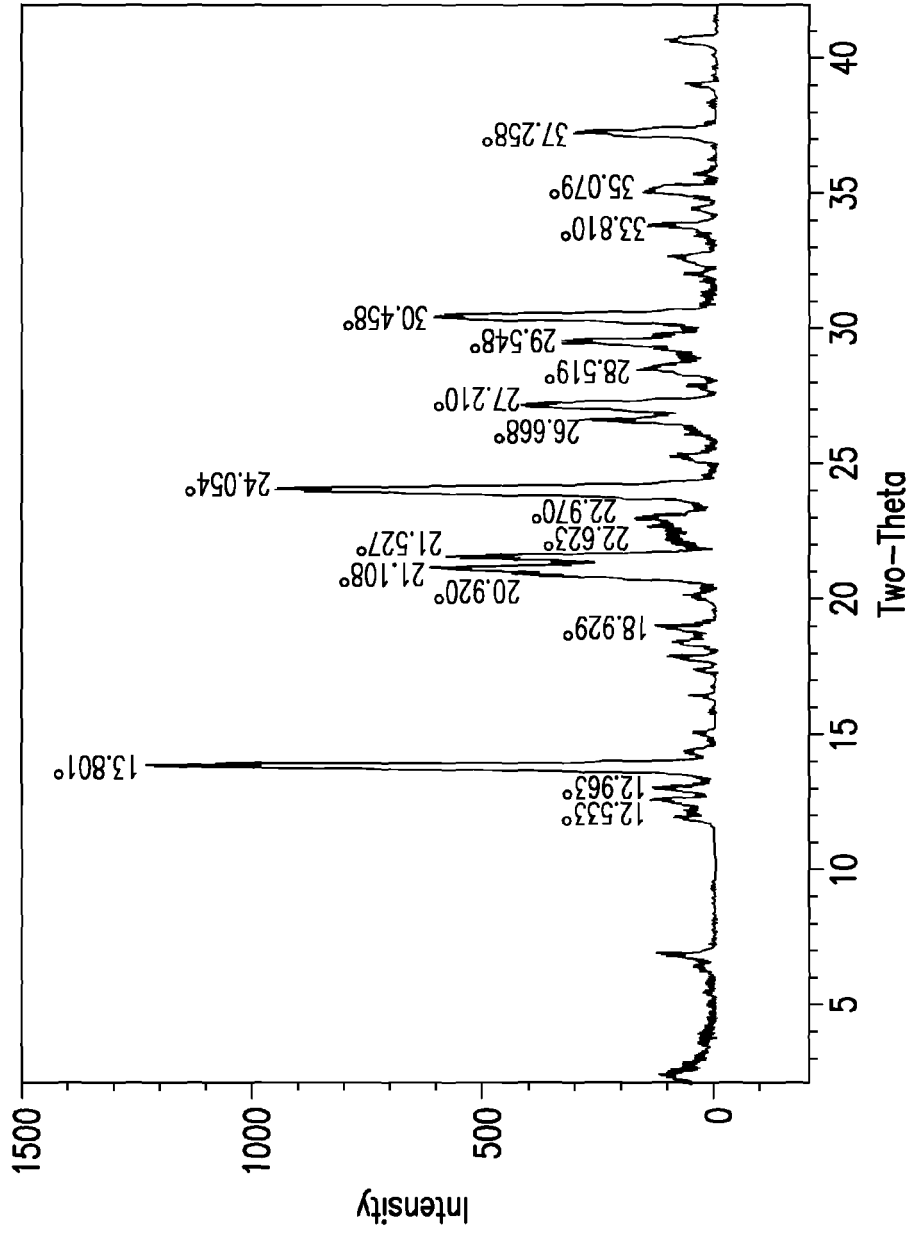


FIG.6

X-ray Powder Diffraction Pattern of Azacitidine, Mixed Phase Forms I and VII, Labeled with the more Prominent 2θ Angles (Cu Kα Radiation)

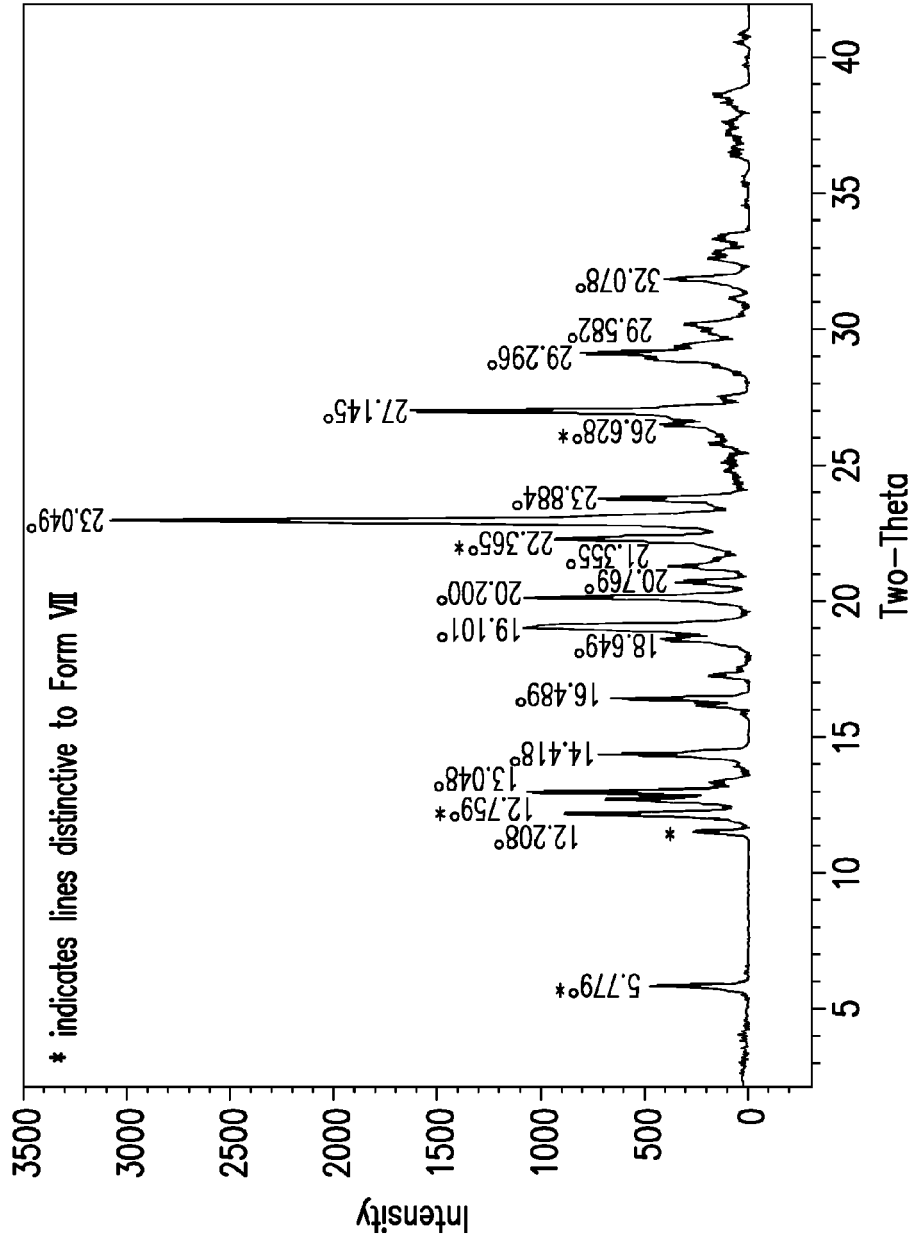


FIG.7

X-ray Powder Diffraction Pattern of Azacidine, Form VIII, Labeled with the more Prominent 2θ Angles (Cu Kα Radiation)

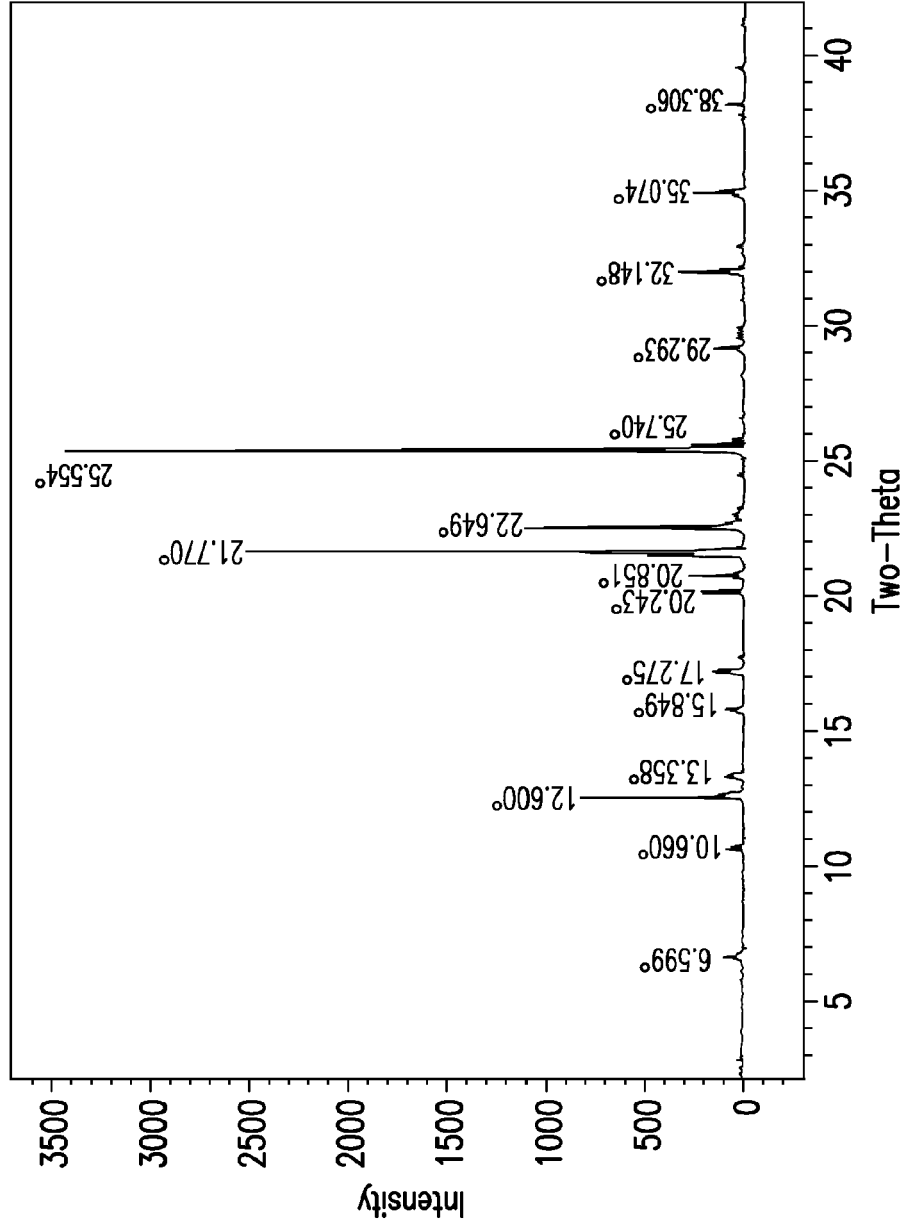


FIG.8

REFERENCES CITED IN THE DESCRIPTION

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