

Case study

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2 Novel Therapy May Be the First Line Treatment for Multiple Myeloma but Should Not Be
3 the Last Word: Two Cases Illustrated

4 ABSTRACT

5 Over the past 20 years, the treatment for multiple myeloma (MM) has evolved
6 significantly. These pharmaceutical developments allow physicians to combine existing
7 chemotherapy with newly approved novel and targeted medications to create various
8 treatment regimens for MM. These novel drug combinations, immunomodulatory drugs
9 (Thalidomide, lenalidomide and Pomalidomide) and proteasome inhibitors (Bortezomib
10 and carfilzomib), are used upfront for induction therapy as well as for maintenance and
11 treatment of subsequent relapses. However, the emergence of resistant myeloma
12 clones to these drugs is usually inevitable. We describe 2 cases here that demonstrate
13 beneficial response to old traditional chemotherapy combinations after patients become
14 resistant to all novel drugs available. Therefore, our main message is that while novel
15 drugs should be used in frontline combinations to treat MM patients, these novel drugs
16 should not be the last word, and often going back to the old traditional chemotherapy
17 may illicit response and possibly prolong survival.

18 INTRODUCTION

19 Over the past 20 years, the treatment for multiple myeloma (MM) has evolved
20 significantly. Between 1995 and 2015, ten new drugs were approved by the FDA for the
21 treatment of MM. These pharmaceutical developments allow physicians to combine
22 existing chemotherapy with newly approved medications to create various treatment

23 regimens for MM. These treatment regimens, when combined with autologous stem cell
24 transplantation (ASCT), play an important role in significantly extending MM patients'
25 survival (1). However, the innumerable amount of different treatment regimens does not
26 allow for comprehensive comparisons of efficacy using phase I-III clinical trials.

27 In most patients, the natural history of MM includes recurrent relapses and death
28 from resistant disease despite these new treatment options. Relapses in MM often
29 occur after treating a heterogeneous malignant plasma cell population due to the
30 emergence of resistant clones (2). Our two cases illustrate the story of two patients,
31 young and old, with MM who received several different treatment regimens. The
32 treatments included both evidence-based regimens and non-evidence based drug
33 combinations, which delayed the emergence of a resistant clone. These two cases also
34 raise ethical considerations regarding access to care, treatment cost, and timing of
35 palliative care and hospice intervention.

36 **Case 1:** A 39-year-old Caucasian female presented at the age of 28 year old, 34 weeks
37 pregnant, presented with swelling and numbness in June 2004 with severe pain while
38 chewing food and was unable to open her mouth. CT scan showed a large mass of
39 ~~starting from~~ the ramus of the left mandible with soft tissue extension and involvement
40 of the left inferior alveolar nerve and evidence of pathological fracture. A biopsy of the
41 mass revealed a plasmacytoma. Further evaluation included a bone marrow (BM)
42 biopsy which showed 10% CD138+ plasma cells. Cytogenetics analysis was normal,
43 while skeletal survey was normal except for the aforementioned mandibular lesion. She
44 was treated with radiation therapy (RT), total of 4000 cGy, followed by observation (in
45 another practice). About 8 months later, she presented with hypercalcemia and back

46 pain. She The patient was diagnosed with progressive multiple myeloma IgG kappa,
47 stage IIIA (stage II by International Staging System [ISS]), causing pathological
48 fractures in T9, left anterior superior iliac spine and right inferior pubic ramus. Bone
49 marrow biopsy showed 30-50% abnormal plasma cells, normal cytogenetics analysis,
50 and monosomy 13 in 5% revealed by FISH. She was treated with 2 cycles of VAD (3)
51 with minimal response. She The patient was subsequently treated with one cycle of
52 HyperCVAD part A (4), followed by peripheral blood stem cell collection and first
53 autologous stem cell transplantation (ASCT) using conditioning regimen of melphalan
54 200 mg/m² (in middle of 2005). Three months post ASCT, repeat evaluation revealed
55 complete morphologic and molecular remission (according to the International Myeloma
56 Working Group criteria) (5) and patient started maintenance on phase II study using
57 interferon alpha (IFN) 4 million units and GM-CSF 125 mcg/m² both given
58 subcutaneously (SC) 3 times weekly (6). Patient became pregnant while on IFN and
59 developed relapse manifesting with hair line fracture of her left tibia which was treated
60 with RT 2500 cGy. She had natural delivery of healthy baby in middle of 2007.
61 Meanwhile, she developed a left distal humerus plasmacytoma eroding the bone cortex
62 and she underwent prophylactic internal fixation. Following that, she patient was started
63 on lenalidomide (Len) and weekly dexamethasone (dexa) (Rd) (7), but had only stable
64 disease and oral cyclophosphamide (Cy) 500 mg given weekly was added. She
65 achieved partial response with < 5% residual plasma cells on repeat BM biopsy, but
66 cytogenetics showed for the first time cell population with hyperdiploidy 56, XX in 3/30
67 metaphases and 2/30 metaphases had del 20q11.2.

68 In 2008, about 3 years after her 1st ASCT, she the patient had a second ASCT with
69 high-dose melphalan 200 mg/m². She achieved VGPR with < 5% plasma cells in the
70 marrow and residual elevation of kappa at 5.05 mg/dL (normal range 0.33-1.94 mg/dL).
71 Her cytogenetics showed 1/30 metaphases with hyperdiploidy 55, XX and del 17 and
72 18. After second transplant, the patient was on oral Cy maintenance 200 mg daily for 10
73 days every 4-6 wks for about 2 years. She remained stable until Aug 2010 when she
74 developed chemical progression. She Patient was started on Doxil, bortezomib (Bor,
75 Velcade) and dexamethasone (Dexa) (8) for 3 cycles and then stayed on maintenance Bor (1.3
76 mg/m² SC weekly for two wks on and 1 wk off) for another 4 months when she showed
77 chemical signs of progression and developed worsening neuropathy with pain in her
78 legs. She was switched to a new regimen consisting of IV Cy 750 mg/m² and liposomal
79 doxorubicin (Doxil) 30 mg/m² for one cycle and for the 2nd cycle oral etoposide 100 mg
80 daily for 5 days was added, each cycle was given every 3 wks and continued for total of
81 13 cycles. The main side effect of this regimen was grade 2/3 mucositis. Chemical
82 progression was diagnosed again in Feb 2012, and at this time, she was treated with
83 subcutaneous Bor weekly, oral Cy 100 mg daily and dexamethasone 20 mg weekly without
84 response. At this point, she the patient was admitted and given one cycle of hyperCVAD
85 part A without significant response. Therefore, her treatment was switched to VTD-
86 PACE (9) given in the inpatient setting for two cycles and with good response achieving
87 VGPR. She was placed on VTD maintenance for 5 months. In Dec 2012, she showed
88 signs of progression and was started on carfilzomib (Carf) single drug at the
89 recommended doses per the manufacturer (Onyx Pharmaceuticals, Inc.). She Patient
90 had significant incremental elevation of liver enzymes after each cycle and the

91 treatment was discontinued during the 3rd cycle, however, she responded and achieved
92 80% reduction in her serum free light chain which plateaued at around 13 mg/dL.
93 Because of the liver toxicity, she was switched to pomalidomide (Pom) 4 mg daily for 21
94 days every 4 wks with no response and therefore added oral Cy 200 mg daily and
95 prednisone 80 mg daily on days 1-5 for each subsequent cycle. Five months later,
96 markers were increasing, and she was switched to Len/Carf/dexa (See Table 1) every
97 28 days. This time, her liver function tests remained normal. While recovering from the
98 3rd cycle in Dec 2013, shepatient developed severe neck pain and was diagnosed with
99 a new destructive lesion in C2. She had neck brace and received 2000 cGy of RT with
100 good clinical response. During that time, her kappa was up to 142.4 mg/dL and she was
101 started on thalidomide (Thal) 100 mg daily and weekly dexa, and then Carf was added
102 after RT was completed at 20 mg/m² days 1,2,8,9,15,16 every 4 wks for one cycle with
103 progressive disease. She then received VBMCP (10) regimen in the outpatient setting
104 for one cycle without response. Vorinostat was added at 200 mg daily orally for 5 days
105 every week for the second cycle. Meanwhile, pain and swelling developed in her
106 previously involved left humerus. Imaging showed progression with extension of her
107 myeloma into the soft tissues. She received RT, 2000 cGy in 10 fractions, with excellent
108 response. At this point, she the patient was started on weekly SC Bor 1.6 mg/m² with
109 vorinostat 100 mg daily (which was increased to 400 mg daily in the 2nd cycle), dexa 40
110 mg weekly, repeated every 3 wks. Because of lack of response, Thal 100 mg daily was
111 added and then switched to full dose Carf with weekly dexa, daily Thal 100 mg, and
112 vorinostat for one cycle. Now 10 and 1/2 years from diagnosis, with lack of response to
113 the Carf combination, she was started on DT-PACE. She Patient had good response,

114 both symptomatically (improved bone pain) and chemically (drop in her kappa to as low
115 as 12.31 mg/dL). The regimen was given during 6-days hospitalization every 4-5 wks
116 with pegfilgrastim (neulasta). The patient developed pancytopenia after each cycle and
117 required hospitalization on 2 occasions for neutropenic fever and septicemia, including
118 a brief trip to the intensive care unit. After the 3rd cycle, she was able to go with her 5
119 children and husband on an organized trip. SheThe patient was given oral Thal 100 mg
120 daily and SC interferon-alpha (3×10^6 units three times weekly) maintenance regimen.
121 Upon her return, she showed chemical progression again and was admitted for another
122 cycle of DT-PACE. She received her last cycle in Feb 2015 which was complicated by
123 pancytopenia and bilateral pneumonia from which she recovered. The patient course
124 from diagnosis to the end of 2014 is illustrated in Fig 1 with more than 7 chemical and
125 clinical relapses. During all that time, she was actively taking care of the household and
126 her five children with estimated Karnofsky score of 70%. SheThe patient had help and
127 support from her parents and one of them always came with her each clinic visit. Over
128 the last year of her life, she had developed significant muscle wasting and weight loss,
129 likely from high myeloma tumor mass. Patient eventually died from her progressive
130 disease and bilateral pneumonia almost 11 years after diagnosis. (See Table 1 for
131 details of regimens used).

132

133 **Case 2:** An 80-year-old Caucasian male who was diagnosed with kappa light chain
134 multiple myeloma stage IIIB (stage III by ISS) at the age of 74 years. He presented with
135 anemia and fatigue and found to have acute renal failure with creatinine of 4 mg/dL.
136 Skeletal survey showed one lesion in vertebrae L3, and cytogenetic analysis showed

137 normal male karyotype but FISH was positive for Del 13 and IgH gene locus
138 rearrangement. He was started on combination of Cy, Bor and dexamethasone (CVD) (11) for 3
139 cycles, then proceeded to have high-dose melphalan 140 mg /m² and ASCT. He The
140 patient achieved complete remission with improved creatinine to baseline of about 1.4
141 mg/dL. He was placed on oral Cy maintenance 200 mg/day X 10 days every month for
142 18 months, he eventually had chemical progression/relapse, and dexamethasone 40 mg X 4 days
143 every 2 wks was added for one cycle, followed by CVD with minimal response. He was
144 switched to Len 15 mg (dose adjusted according to kidney function) and low dose dexamethasone
145 (7) and had that for 14 months, then Bor was added for 2 more cycles due to disease
146 progression. Due to minimal response, hepatient was started on a single agent Carf and
147 received 15 cycles before showing signs of laboratory progression. He received one
148 cycle of CVD without response, then changed treatment to Pom 4 mg/day for 21 days
149 every 28 days and dexamethasone 40 mg weekly for 6 months until he stopped responding. At this
150 time, he was started on our modified VBMCP, which is given without Mel (VBCP, see
151 Table 1) all in the outpatient clinic, followed by pegfilgrastim. So far, patient has
152 received 8 cycles of VBCP, with continuous response achieving partial remission. He is
153 supported with blood transfusions due to persistent thrombocytopenia. Again, charting
154 his kappa light chain levels over the years (Fig 2) demonstrates lack of CR with shorter
155 responses in the last 2.5 years. His last bone marrow biopsy was done about 14 months
156 ago and showed 40% plasma cells by CD138 immunohistochemistry, while his
157 cytogenetics showed 48 X,-Y and complex abnormalities in 12 metaphases including
158 t(8;14), t(1;8), t(3;13), del 13, trisomies 11, 15, 19 and 21, as well as del 16 and 20.
159 FISH studies showed del 13, IGH/MYC gene loci fusion, and amplification of MAF gene.

160 One may question quality of life considering that he is in clinic twice weekly, with the
161 great support of his wife. HeThe patient was wheel chair bound due to muscle wasting
162 and undergoes home physical therapy, with Karnofsky score of 50%. However, he
163 otherwise enjoyed his daily activities, including reading, listening to his favorite music,
164 and entertaining his friends in the comfort of his home. (See Table 1 for details of
165 regimens used)

166 DISCUSSION

167 Our two cases illustrate several important points in the treatment of MM. In this report
168 we demonstrated the use of different combinations in sequential and continuous
169 manner to keep the patient with MM alive. It seems that the natural history of recurrent
170 resistant relapses has not changed but rather delayed and stretched because of the
171 various available and effective therapies. Furthermore, the treatment options used in
172 these two cases may not be available for many myeloma patients in other Western and
173 developing countries. Even in our own community, a patient like the ones presented
174 here may have been referred to palliative care/Hospice earlier in the course of their
175 disease, which may have lead earlier death. From that point of view, these cases may
176 not be an unusual case of MM, but rather represents other similar cases with similar
177 disease course, even in older patients, receiving this kind of sequential therapy. It is
178 also important to think about these cases in view of the discussions in regards to the
179 cost of all the new novel drugs and access to care. Our patients, had
180 Medicaid/Medicare or just Medicare alone, that covered all the treatments including two
181 ASCTs in case 1. However, allogeneic transplant, which some transplant experts
182 consider the best next step for young MM patients like case 1, was not possible

183 because it is not covered by Medicare. Considering that case 1 was a young mother of
184 5 children, then we believe that the cost per year saved may have been fully justified.

185 Few points emerged from these teaching cases regarding the use of the
186 available drug combinations: 1. Both cases illustrate the use of same drugs again (such
187 as Cy, Bor, dexamethasone and ASCT) in repeat cycles and at later time in the course of the
188 disease with positive effect; 2. One can mix and match drugs at different time points in
189 order to elicit more responses, albeit without existing evidence in the literature; 3. Old
190 chemotherapy drug combinations, with or without novel agents, can work even when
191 newer novel drugs stop working. Furthermore, in case #2, the VBCP served as a bridge
192 for newer potentially less toxic therapies that were approved recently in 2015.

193 CONCLUSION

194 Thus our conclusion is that while novel drugs should be used in frontline
195 combinations to treat MM patients, these novel drugs should not be the last word, and
196 often going back to the old traditional chemotherapy may illicit response and possibly
197 prolong survival.

198 **Figure Legends**

199 **Fig 1:** The illustration of the time course (2004-2015) of case 1 with relapses
200 documented in the bottom panel showing the curve for changes in serum free kappa
201 light chain (mg/dL, Y axis) generated by EPIC medical records and demonstrating that
202 with time (X axis), remissions became shorter and the relapses (indicated by spikes in
203 kappa light chain) got more frequent and more resistant as illustrated (upper panel, Y
204 axis) by the number of different regimens used. (**Abbreviations:** ASCT, autologous

205 stem cell transplantation; IFN, interferon alpha; RT, radiation therapy; Lt, left; C2,
206 cervical vertebra 2; T, thoracic vertebra).

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208 **Fig 2:** Time course of Case 2 (2009-2016, yellow colored bar) which shows the curves
209 for kappa (mg/dL, Y axis) and Kappa/lambda ratio as well as the different treatment
210 given at times of disease progression. The lettered lines represent the length of different
211 treatments given to the patient after ASCT. The X axis shows the time line for his light
212 chain measurements and the different treatment given. A, Cy maintenance. B, Cy
213 +dexa. C, CVD. D, Len +dexa with and without Bor. E, Carf as single agent. F, Carf +Cy
214 (500 mg weekly) +dexa. G, Pom +dexa. H, Pom +Bor +dexa. K, VBCP, so far 8 cycles.
215 (**Abbreviations:** Cy, cyclophosphamide; dexa, dexamethasone; CVD, Cy +Velcade
216 [bortezomib] +dexa; Len, lenalidomide; Carf, carfilzomib; Pom, pomalidomide; Bor,
217 bortezomib; VBCP, vincristine +BCNU [carmustine] +Cy +Prednisone).

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257 Table 1: Details of chemotherapy regimens used in these two cases.

Regimen	Schedule of drug delivery	Frequency	Growth factor Yes/No
HyperCVAD* (inpatient)	Days 1-3: Cy 300mg/m ² IV every 12 hr (with mesna) Days 4-5: Doxorubicin 50mg/m ² & vincristine 2mg given in continuous IV for 48 hrs Days 1-5: Dexamethasone 40mg orally	Every 28-35 days	Yes
DVD (Outpatient)	Days 1, 4, 8, and 11: Bor (V) 1.3mg/m ² SC Day 4: Pegylated liposomal doxorubicin 30mg/m ² IV over 60 minutes. Days 1-4: Dexamethasone 40mg orally	Every 3 wks	No
VTD-PACE** (Inpatient)	Days 1, 4, 8, and 11: Bor 1 mg/m ² SC Continuous: Thalidomide 50–200mg orally daily at bedtime Days 1-4: Dexamethasone 40mg orally daily Days 1-4: Cyclophosphamide 400mg/m ² + etoposide 40mg/m ² + cisplatin 10mg/m ² + doxorubicin 10mg/m ² , all given in continuous IV infusion over 24 hours daily.	Every 28-35 days	Yes
Carf/Len/Dexa	Carf 20 mg/m ² IV on days 1,2,15, 16; Len 25 mg daily PO X 21 days; dexa 40 mg weekly.	Every 28 days	No
VBMCP (outpatient)	Day 1: vincristine 1.2mg/m ² (limit of 2 mg) IV, BCNU 20mg/m ² IV, Cy 400mg/m ² IV Days 1-4: oral Mel 8mg/m ² Days 1-7: oral prednisone 40mg/m ²	Every 35 days	Yes/No
VBCP*** (Outpatient)	Day 1: vincristine 1.2mg/m ² (limit of 2 mg) IV, BCNU 20mg/m ² IV Days 1-4: Cy 400mg/m ² IV (± mesna) Days 1-7: oral prednisone 40mg/m ²	Every 28 days	Yes

CVD (outpatient)	Days 1,8,15: Cy 500mg oral or IV Days 1,4,8,11: Bor 1.3mg/m ² SC and Dexa 40mg [†] on the day of and the day after Bor.	Every 21 days	No
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258 **Abbreviations:** Bor. Bortezomib; Dexa, dexamethasone; Cy, cyclophosphamide; Mel,
259 melphalan.

260 *Day 11 of this regimen was not given in our practice.

261 **DT-PACE is the same regimen but without Bor (Velcade). Usually days 8 and 11 are
262 omitted in our practice.

263 ***This regimen was developed in our practice, separate manuscript on our experience
264 is in preparation.

265 † After first cycle, dose is reduced to 20 mg.

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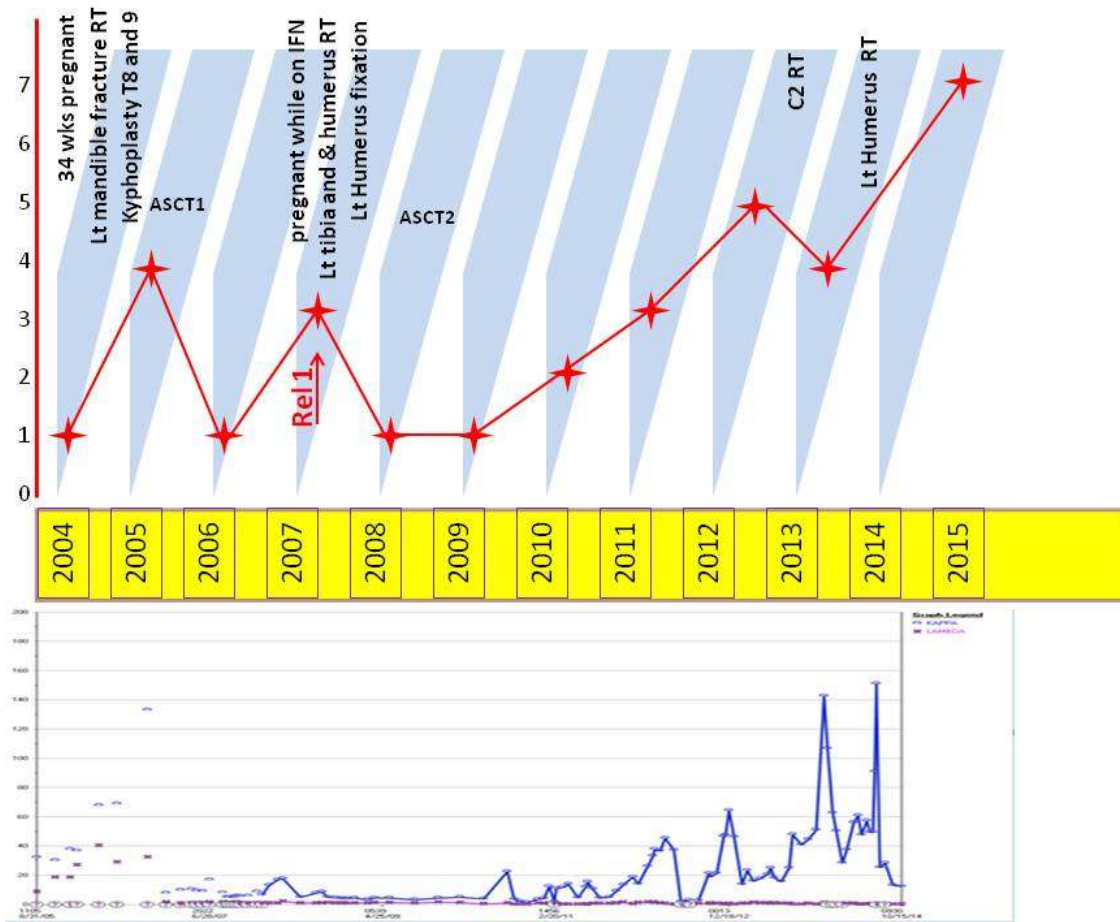
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276 Figure 1.



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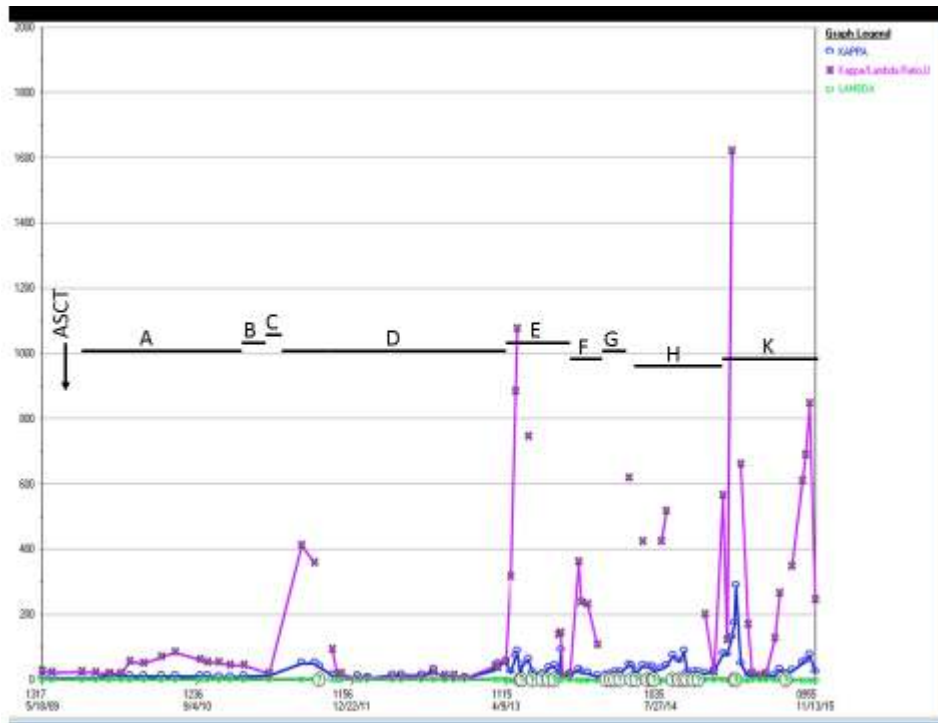
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284 Figure 2.



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