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3 Original Research Article

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5 BREAKTHROUGH CANCER PAIN: PRELIMINARY DATA OF THE Italian Oncologic
6 Pain multiSetting Multicentric Survey (IOPS-MS)

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82 **ABSTRACT**

83 **Introduction:** An ongoing national multicenter survey (Italian Oncologic Pain multiSetting
84 Multicentric Survey [IOPS-MS]) is evaluating the characteristics of breakthrough cancer pain
85 (BTP) in different clinical settings. Preliminary data from the first 1500 cancer patients with
86 BTP enrolled in this study are presented here.

87 **Methods:** Thirty-two clinical centers are involved in the survey. A diagnosis of BTP was
88 performed by a standard algorithm. Epidemiological data, Karnofsky index, stage of disease,
89 presence and sites of metastases, ongoing oncologic treatment, and characteristics of
90 background pain and BTP and their treatments were recorded. Background pain and BTP
91 intensity were measured. Patients were also questioned about BTP predictability, BTP onset
92 (≤ 10 minutes or > 10 minutes), BTP duration, background and BTP medications and their
93 doses, time to meaningful pain relief after BTP medication, and satisfaction with BTP
94 medication. The occurrence of adverse reactions was also assessed, as well as mucosal
95 toxicity.

96 **Results:** Background pain was well controlled with opioid treatment (numerical rating scale
97 3.0 ± 1.1). Patients reported 2.5 ± 1.6 BTP episodes/day with a mean intensity of 7.5 ± 1.4 and
98 duration of 43 ± 40 minutes. 977 patients (65.1%) reported non-predictable BTP, and 1076
99 patients (71.7%) reported a rapid onset of BTP (≤ 10 min). Higher patient satisfaction was
100 reported by patients treated with fast onset opioids.

101 **Conclusions:** These preliminary data underline that the standard algorithm used is a valid
102 tool for a proper diagnosis of BTP in cancer patients. Moreover, rapid relief of pain is crucial

103 for patients' satisfaction. The final IOPS-MS data are necessary to understand relationships
104 between BTP characteristics and other clinical variables in oncologic patients.

105 **KEY WORDS**

106 Breakthrough pain, cancer pain, pain assessment, rapid onset opioid.

107 INTRODUCTION

108 Pain is common in cancer patients, particularly in the advanced stage of disease when the
109 prevalence is estimated to be more than 70% [1]. Adequate pain control is achieved in most
110 patients with available analgesic therapies [2]. However, despite adequate pain control for
111 most hours of the day, patients may develop transient flares of pain throughout the day. This
112 phenomenon is known as breakthrough cancer pain (BTP) [3]. BTP has been reported to
113 produce a negative impact on quality of life and is associated with a significant physical,
114 psychological and economic burden [4]. Several studies have assessed the epidemiology of
115 this phenomenon, reporting largely variable data in different settings by using different
116 definitions and methodologies, for example without an *a priori* definition of BTP, without
117 clearly distinguishing background pain intensity and BTP intensity, or without considering
118 the level of opioids used for background analgesia [5-7]. In recent years, BTP has been more
119 meaningfully characterized through a diagnostic algorithm. Moreover, some attempts to
120 better characterize this phenomenon according to a number of variables have been made.
121 Recently, an expert consensus suggested that a BTP subclassification according to the
122 characteristics of BTP may provide tailored treatment [8].

123 In the previous Italian Oncologic Pain multiSetting (IOPS) study, performed in various settings
124 in a large number of patients, several factors influencing the development and characteristics
125 of BTP were assessed [9]. From this data, the IOPS expert group planned a new multicenter
126 survey, with the aim of providing further information on BTP and the factors influencing its
127 characteristics in a large number of patients, diagnosed according to a specific algorithm. The
128 use of BTP medications and factors interfering with administration of transmucosal opioids,
129 commonly used for the management of BTP because their PK profile fits with BTP onset and

130 duration, were also evaluated [5]. Reported here is a preliminary analysis of data from the
131 first 1500 patients of 4056 patients globally enrolled in this second IOPS study.

132 **METHODS**

133 This preliminary analysis included the first 1500 patients recruited in a national,
134 observational, multicenter Italian study. An investigator meeting was held to present and
135 comment on the project with the representatives of each center that participated.

136 Subsequently, each center received an IOPS Multicentric Survey (IOPS-MS) investigator
137 manual.

138 Thirty-two centers were involved. Each center consecutively enrolled patients for 24 months
139 after obtaining local ethic committee approval and the patients' informed consent. Patients
140 were recruited in the most common care settings where cancer patients are assessed for pain,
141 including oncology, outpatient pain therapy, palliative care, and radiotherapy settings. The
142 place of assessment was also recorded, including outpatient clinic, day hospital, home care,
143 hospice, and inpatient ward.

144 Inclusion criteria were: age >18 years, cancer diagnosis at any stage, stable background pain
145 in the last week with an intensity of ≤ 4 on a numerical scale from 0 to 10, and episodes of BTP
146 with an intensity of ≥ 5 , clearly distinguished from background pain. A standard algorithm to
147 diagnose BTP was followed according to the following definition: BTP is a transitory
148 exacerbation of pain of moderate to severe intensity that occurs spontaneously or predictably
149 [8-11], and is well distinguished from background pain of mild intensity [6, 12]. Exclusion
150 criteria were the absence of a cancer diagnosis, uncontrolled background pain (>4 on a
151 numerical scale of 0 to 10), or no relevant increases in pain intensity (<5) which could be

152 interpreted as BTP episodes. Patients unable to provide information about the data required
153 for the study, due to either cognitive failure or terminal disease, were also excluded. Patients
154 meeting the inclusion criteria and assessed at each center were consecutively surveyed.

155 Epidemiological data, Karnofsky index, stage of disease, presence and sites of metastases,
156 ongoing oncologic treatment, and characteristics regarding background pain and BTP and
157 their treatments were recorded. Type of pain was registered according to routine clinical
158 practice (neuropathic, nociceptive or coexistent mechanism), and background and BTP
159 intensity were measured on a numerical scale from 0 to 10. Patients were also questioned
160 about BTP predictability, BTP onset (≤ 10 minutes or > 10 minutes), BTP duration, background
161 and BTP medications and their doses, time to meaningful pain relief after BTP medication, and
162 satisfaction with BTP medication (a 4 point scale was used by physicians: very satisfied,
163 satisfied, not satisfied, and neither satisfied nor dissatisfied) [9,10]. The occurrence of adverse
164 reactions was also assessed, and mucosal toxicity was graded according to the World Health
165 Organization (WHO) criteria [13]. The presence of candidiasis and xerostomia was also
166 recorded. Each patient followed local policy and therapeutic protocols, and no specific
167 treatment for BTP was assigned. To guarantee good quality of the data, these were entered in
168 a web-based clinical report form. Each center had an individual password to enter their data
169 into the system, and the study monitors could check records by local and remote monitoring.

170 **Statistics**

171 Data from the first 1500 patients were preliminarily examined. Continuous variables were
172 summarized as means and standard deviations (SD). Categorical variables were summarized
173 as percentages (absolute numbers). Univariate analysis was performed using the Wilcoxon or
174 chi-square test without correction for continuity for comparison among groups of continuous

175 and categorical variables, respectively. Multivariate analysis was based on generalized linear
176 models, with suitable link function chosen according to the characteristics of the response
177 variable: identity for continuous and logit for binary or proportional-odds ordered categorical
178 variables. All variables considered were entered into the model as they were, without any
179 transformation or cut-off. The nonlinear effect of covariates was modeled by means of a
180 restrictive cubic spline function, and its significance was assessed by means of the χ^2 Wald
181 test. The model strategy was determined by following a backward selection strategy among
182 variables reaching a level of at least 0.25 on univariate analysis. Model fit was considered
183 significantly improved on the basis of the Akaike information criterion (AIC) applied
184 backward for each model at a significance level of 0.05. To avoid inflation in type-I error due
185 to multiplicity of testing, subgroup analysis was conducted by introducing interaction terms
186 into the main multivariate model, and its significance assessed by means of AIC. Multivariate
187 models were depicted as nomograms. To evaluate the goodness of fit of the models, cross-
188 validation and bootstrap (1000 runs) techniques were applied by the use of Somer's D_{xy} .
189 Statistical significance was set at $p \leq 0.05$. The R-System statistical package and the Harrell
190 regression modelling strategies libraries were used for analysis.

191 **RESULTS**

192 **Patient characteristics**

193 Of the first 1500 patients recruited in IOPS-MS, most had metastatic disease and were
194 receiving anticancer treatment (Table 1). The most common care settings were oncology and
195 pain therapy, and patients were seen most often in outpatient clinics (37%) and inpatient
196 wards (33%). No differences in gender were found among the different settings ($p=0.989$). A
197 lower and a higher Karnofsky index were found in the palliative care and radiotherapy

198 settings, respectively (39.4 ± 10.8 vs 70 ± 18.2 ; $F=86.7$; degrees of freedom [d.f.]=3.519;
199 $p<0.001$). Finally, older patients (mean \pm SD age 73.9 ± 12.5 years) were over-represented in
200 the palliative care setting ($F=27.1$; d.f.=3.519; $p<0.001$).

201 **BTP characteristics**

202 The initial diagnosis of BTP was most often performed by oncologists ($n=616$ diagnoses, 41%)
203 and pain physicians ($n=583$, 39%), followed by palliative care physicians ($n=241$, 16%),
204 nurses ($n=18$, 1%), general practitioners ($n=15$, 1%), other healthcare providers ($n=15$, 1%),
205 and radiotherapists ($n=9$, 0.6%). In three cases, data were unavailable. Patients in hospices
206 had a longer time from diagnosis of BTP in comparison with outpatient settings ($p=0.0123$).
207 The percentages of patients with baseline pain and the characteristics of BTP are presented in
208 Figure 1.

209 The mean number of BTP episodes/day was 2.5 ± 1.6 (data available for 1499 patients). In
210 patients with higher Karnofsky index and with prostate cancer the number of BTP episodes was
211 significantly higher than in patients with other primary diagnoses ($p<0.001$). BTP onset was
212 ≤ 10 minutes and >10 minutes in 1076 (71.7%) and 424 (28.3%) patients, respectively.

213 The mean duration of untreated BTP was 43 ± 40 minutes (data available for 504 patients).
214 Variables significantly associated with a longer BTP duration were metastatic disease
215 ($p=0.03$), head and neck cancer and pancreatic cancer ($p=0.04$), and receiving anticancer
216 therapy ($p=0.05$; Table 2). In the multivariate analysis, a significant association with
217 background pain intensity was found, with a linear effect of 10.9 min (95% confidence
218 interval [CI] 9.3 to 12.5).

219 The distribution of BTP mechanisms in the different care settings is reported in Table 3. A
220 mixed mechanism of BTP was found to be more represented in oncology and pain therapy
221 settings than in radiotherapy and palliative care settings. Conversely, a nociceptive
222 mechanism was more frequently found in palliative care and radiotherapy settings than in
223 oncology and pain therapy settings.

224 **Predictable BTP**

225 BTP was unpredictable in 977 patients (65.1%) and predictable in 523 patients (34.9%).
226 Predictable BTP was associated with age ($p=0.008$), pain mechanism ($p<0.001$, lower risk
227 with mixed mechanism), place of assessment ($p<0.001$), care setting ($p=0.002$), background
228 pain ($p=0.004$), diagnosis of prostate cancer ($p=0.030$), Karnofsky index ($p=0.046$) and oral
229 mucositis ($p<0.001$). In the multivariate analysis, lower Karnofsky, lower BTP intensity, and
230 rapid onset of BTP were significantly associated with predictable BTP. The radiotherapy
231 setting was strongly associated with predictable BTP (odds ratio [OR] 9.05). The main trigger
232 for predictable BTP was activity-movement ($n=349$, 67%), followed by swallowing ($n=80$,
233 15%), cough ($n=54$, 10%), procedure ($n=39$, 7%), and bowel movement ($n=31$, 6%).

234 **Intensity of background pain and BTP**

235 The mean intensity of background pain on assessment and the average pain in the previous
236 week were 3.0 ± 1.1 , and 3.0 ± 1.1 , respectively. The mean doses of oral morphine equivalents
237 (OME) used for background pain were 69.8 ± 139.7 mg/day. The mean intensity of BTP was
238 7.5 ± 1.4 . Rapid-onset BTP and high levels of background pain intensity were associated with
239 a higher BTP intensity. Conversely, a slow-onset BTP was associated with a lower BTP
240 intensity. No differences in BTP intensity among the care settings and triggers of predictable

241 BTP were found. Using mixed pain mechanism as a reference, BTP intensity was higher for
242 neuropathic pain ($p=0.0248$) and lower for nociceptive pain ($p=0.0257$). BTP was of lower
243 intensity in older patients ($p=0.0002$), in patients with higher Karnofsky status ($p=0.0016$),
244 and in patients with breast cancer ($p=0.04$). Finally, mucositis was associated with higher BTP
245 intensity ($p=0.0083$).

246 **BTP medications**

247 A total of 1263 (84%) patients were receiving opioid drugs for the management of BTP,
248 including fentanyl pectin nasal spray (FPNS, 23%), oral morphine (OM, 17%), fentanyl buccal
249 sublingual tablet (FBST, 15%), fentanyl buccal tablet (FBT, 11%), oral transmucosal fentanyl
250 citrate (OTFC, 5%), subcutaneous morphine (SC-M, 4%), intravenous morphine (IV-M, 3%),
251 and intranasal fentanyl spray (INFS, 1%). The mean \pm SD doses of each drug were FPNS (178
252 ± 144 μg), OM (13 ± 11 mg), FBST (227 ± 169 μg), FBT (261 ± 207 μg), OTFC (490 ± 330 μg),
253 SC-M (11 ± 5 mg), IV-M (9 ± 9 mg), and INFS (109 ± 59 μg). No differences in BTP medication
254 according to the characteristics of BTP were found. FPNS was less frequently used in
255 radiotherapy and pain therapy settings ($p=0.008$), while SC-M was more frequently used in
256 oncology and palliative care settings ($p=0.004$). There was a significant relationship between
257 OME and opioid doses for BTP (correlation 0.42, 95% CI 0.37 to 0.46).

258 **Time to meaningful pain relief after drug administration**

259 The mean time for achieving meaningful pain relief after BTP medication was 17 ± 14
260 minutes. In Table 4, the variables associated with the time for meaningful pain relief are
261 presented (data were available for 810 patients). In the multivariate analysis, factors
262 associated with shorter meaningful pain relief were assessment in the inpatient ward

263 (p<0.001), drug therapy (INFS, FPNS and IV-M, p=0.012), and pancreas and head and neck
264 cancers (p=0.0193).

265 **Satisfaction with BTP medication**

266 Patients were very satisfied, satisfied, not satisfied, and neither satisfied nor dissatisfied with
267 their BTP medication in 154 (11%), 765 (55%), 262 (19%) and 211 (15%) cases (data
268 available in 1392 patients). The level of satisfaction was significantly associated with the use
269 of FPNS (p=0.0002). Also, the outpatient clinic (p=0.04), care in the oncology setting
270 (p=0.0011), and receiving anticancer treatment (p=0.0166) were associated with higher
271 patients' satisfaction (Table 5).

272 **Adverse effects of BTP medications**

273 Adverse reactions attributed to BTP medications were reported in 53 out of 1500 (4%)
274 patients and were: constipation (n=18), dizziness (n=18), nausea (n=5), headache (n=2),
275 vomiting (n=1) and other unspecified adverse effects (n=9). The intensity was mild in 46
276 patients (88%) and moderate in 6 patients (12%). In 38 patients (83%) no specific
277 therapeutic change was required, while in the remaining 8 cases (17%) it was deemed
278 necessary to treat the adverse effects or discontinue the BTP medication. No association was
279 found between adverse reactions and choice and dosage of opioids used for BTP (p=0.843).
280 Finally, no medication abuse was reported.

281 **Oral mucositis**

282 Two hundred and twelve patients (14%) presented with different levels of oral mucositis. Of
283 them, 134 patients had oral aching/erythema, 56 had oral erythema/ulcer/solid diet

284 tolerated, 17 patients had oral ulcers/only liquid diet tolerated, and in 5 patients oral feeding
285 was impossible (from level 1 to level 4, respectively). Head and neck cancer was positively
286 associated with the severity of oral mucositis (OR 5.42; 95% CI 2.70 to 10.86; $p < 0.001$). Of
287 interest, the grade of mucositis was positively associated with BTP on swallowing (OR 4.85;
288 95% CI 2.79 to 8.40). No association was found between levels of oral mucositis and choice of
289 drugs for BTP and their doses. Candidiasis and xerostomia were detected in 90 (6%) and 280
290 (19%) patients, respectively.

291 **DISCUSSION**

292 Preliminary data for the first 1500 patients of the IOPS-MS survey suggest that, in general, in
293 patients with BTP, older patients and patients with a lower Karnofsky index were most
294 frequently followed in a palliative care setting. This information is consistent with data
295 collected in the previous IOPS survey [9] and in other surveys performed either in oncology or
296 in palliative care settings [14, 15], confirming that the patients' characteristics differ among
297 the settings of care, particularly in patients with the highest morbidity under the care of
298 palliative care physicians. Data suggest that higher prevalence rates of BTP are reported in
299 studies performed in the hospice setting [9, 16, 17].

300 Results of this survey suggest that the diagnosis of BTP was performed more frequently by
301 oncologists than by palliative care physicians. Conversely, a longer time for diagnosis of BTP
302 was reported in the hospice setting. Oncologists generally have more opportunities to make
303 an early diagnosis of BTP, as they see patients more often through the course of disease [18],
304 whereas physicians in palliative care see patients later in the course of their disease, which
305 may explain this result. Another explanation could be that oncologists have improved their
306 pain assessment skills in the years since large surveys showed worrying data, suggesting a

307 great need for continuing education programs in pain management among oncologists [19,
308 20]. However, it is important to note that these findings may not adequately represent the
309 situation, particularly as the differences in the amount of patients with BTP in oncology
310 versus palliative care setting may simply be due to the sampling design. Further investigation
311 is warranted.

312 In this preliminary survey, prostate cancer, a tumor commonly associated with multiple bone
313 metastases, significantly produced more episodes of BTP, potentially representing a risk
314 factor for this phenomenon (see below, predictable BTP). This observation should be
315 confirmed by the complete analysis of the IOPS-MS data. In a European survey, patients had a
316 median of three BTP episodes/day. Of interest, patients were included whether they had just
317 one episode/month or up to 24 episodes/day [10]. Patients who had a better Karnofsky index
318 were more likely to have more BTP episodes. It is likely that more physical activity may
319 produce more episodes of BTP. Alternately, one can argue that the management of
320 background pain of these patients could be better optimized. This observation confirms
321 previous data, in which very advanced and bedridden patients had fewer BTP episodes with
322 longer onset [9].

323 The mean duration of untreated BTP was about 40 minutes, reflecting data from many
324 epidemiological studies that describe a variable duration of 30–60 minutes [9, 10, 21]. BTP
325 duration has been reported to be longer in spontaneous unpredictable BTP than in patients
326 with incident-type BTP [10]. It should be considered that BTP duration in untreated BTP is
327 more difficult for patients to properly assess, and not all patients are able to do so.

328 To facilitate the patients' orientation, a dichotomous measure was chosen for BTP onset (≤ 10
329 or > 10 minutes). BTP onset was rapid in 71.7% of patients and slower in 28.3% of patients.

330 Similar values, with a median of 10 minutes, have been found in a multicenter European
331 survey[10] and an Italian survey[9], where they were lower with incident-type BTP].

332 BTP predictability is an important clinical factor, having obvious therapeutic consequences
333 for timing and choice of available BTP medications. Moreover, incident-predictable BTP has
334 been considered to be a negative factor for cancer pain management [17, 22, 23]. This is due
335 to the difficulties in balancing analgesia at rest and pain on movement, which often results in
336 attempts to improve basal analgesia with a possible occurrence of opioid-induced adverse
337 effects. Predictable BTP has a faster onset, typically observed in patients with bone
338 metastases, triggered by physical activity or movement. In this survey, about 35% of patients
339 had predictable BTP, and physical activity was the most frequent trigger.

340 Some factors were independently associated with predictable BTP and included lower
341 Karnofsky index, lower BTP intensity, and faster BTP onset. Predictable BTP has been
342 previously found to be associated with a faster onset of BTP [9, 10]. Pain induced by
343 movement in patients with bone metastases occurs rapidly and is clearly predictable. A worse
344 performance status was associated with predictable BTP. This is in contrast with a previous
345 finding and probably due to the different care setting distribution in the first IOPS study [9]. It
346 is reasonable to hypothesize that patients with a lower Karnofsky index have lower
347 background pain intensity at rest for most daytime hours, but develop predictable BTP on
348 movement. These data should be confirmed in a larger number of patients with complete
349 analysis the IOPS-MS study. Furthermore, the relationship between predictable BTP and BTP
350 intensity is complex. Patients with a higher BTP intensity had less predictable BTP. This could
351 be explained by patients' attitudes in limiting a sustaining trigger that induced a predictable
352 BTP, thus avoiding a higher peak of pain intensity.

353 Of interest, predictable BTP was more frequently observed in the radiotherapy setting, which
354 could be explained by the fact that patients are commonly referred to these specialists for the
355 treatment of bone metastases.

356 Among the other trigger factors for predictable BTP, swallowing was associated with oral
357 mucositis. Thus, mucosal damage, commonly reported in patients who have received or are
358 still receiving toxic agents [24], is more likely to produce a predictable BTP on swallowing. As
359 expected, mucositis was associated with head and neck cancer, possibly due to previous
360 anticancer treatment. The presence of mucosal damage was also associated with higher levels
361 of BTP intensity. Mucositis is a typical example of BTP occurring with swallowing only.
362 Moreover, the presence of oral mucositis has obvious clinical consequences in terms of route
363 of administration when considering the possible use of transmucosal agents such as rapid-
364 onset opioids, prejudicing reliable absorption of oral transmucosal agents [5]. This suggests
365 that physicians should pay more attention to the diagnosis of mucositis, but also to
366 xerostomia and candidiasis, for optimal selection of BTP therapy.

367 The relationship between background analgesia and BTP intensity is fundamental in
368 describing the phenomenon of BTP, particularly from a therapeutic perspective. It has been
369 reported that a meaningful cut-off of these levels of pain intensity, as reported in the real
370 world by patients instructed in BTP, is about double [12]. In this survey these levels were
371 maintained on average (3 and 7.5 for background pain and BTP intensity, respectively),
372 suggesting that the standard algorithm used in this study allows an appropriate diagnosis of
373 BTP in cancer patients. Of interest, younger patients, higher background pain intensity, a
374 short BTP onset, the level of mucositis, and neuropathic mechanisms were also independently

375 related to BTP pain intensity. These aspects are worthy of further evaluation with the
376 complete data.

377 The relationship between background pain and BTP intensity is problematic. Some patients,
378 for example, avoid taking a medication because BTP intensity is not considered high enough.
379 On the other hand, in a recent Delphi survey, experts in the field of BTP suggested that
380 transient pain exacerbations can occur independently of background pain level and ongoing
381 pain medication, and the phenomenon includes several subgroups of BTP types [8].

382 In our survey, a large number of patients were receiving opioids for the management of BTP,
383 particularly transmucosal fentanyl, in relatively similar or proportional doses, according to
384 the fentanyl availability of different delivery systems. Of interest, a highly significant
385 relationship between the doses of BTP opioid medications and opioid doses for background
386 pain was found. This finding reflects the growing evidence suggesting that a dose proportional
387 to the basal opioid regimen is both safe and effective [25-28], regardless of recommendations
388 suggesting titrating the dose against the effect [29]. Moreover, adverse reactions attributed to
389 BTP medications were limited and of mild intensity in most cases, and were independent of
390 the drug and dose used. This observation confirms that opioid medications given in doses
391 proportional to background opioid dose are relatively safe [25, 26]. This aspect deserves
392 further analysis.

393 Nasal administration of fentanyl provided faster analgesia relative to other fentanyl products
394 [30]. Patient-reported satisfaction with pain treatment is an important outcome measure
395 when assessing both background pain and BTP [8]. Of interest, the use of FPNS and IV-M,
396 home care assessment, pain therapy setting, and the absence of anticancer treatment were
397 associated with the highest level of satisfaction. Therefore, faster analgesia and patients'

398 satisfaction should be strongly considered in order to prescribe optimal treatment. These
399 aspects deserve further research and will be better explored with the complete data of IOPS-
400 MS.

401 There are some limitations to this survey, mainly due to the inherit nature of the study design.
402 Firstly, caution must be taken when interpreting some of the outcomes due to the
403 retrospective nature of the survey. Furthermore, for some outcomes, data are missing.

404 **CONCLUSIONS**

405 Overall, the number of patients allows a preliminary analysis only. Although preliminary,
406 these data provide interesting information that will be developed with the complete IOPS-MS
407 survey. BTP diagnosis was performed according to strict criteria, including stable background
408 analgesia achieved with analgesics given around the clock. BTP intensity was clearly
409 distinguished from basal pain, confirming the validity of the algorithm used for the diagnosis
410 of BTP. These aspects allow us to better evaluate the BTP phenomenon. The characteristics of
411 BTP, including the number of episodes, predictability, onset, intensity, duration, and time
412 from diagnosis were influenced by the many variables taken into consideration. From a
413 therapeutic point of view, opioids, particularly fentanyl products, were largely given for BTP
414 management. The analgesic effect of BTP medications was dependent on a number of
415 variables. Satisfaction with BTP medications was relatively good, particularly in specific
416 settings and with fentanyl preparations. Tolerability was acceptable in most cases,
417 independently of the medication used. Despite the presence of oral mucositis, there was no
418 association with specific drugs or delivery systems. Further data from IOPS-MS should
419 provide a more complete picture of BTP in patients with different cancer types receiving
420 various anticancer treatments, to finally understand this “phenomenon”.

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424 All named authors meet the International Committee of Medical Journal Editors (ICMJE)
425 criteria for authorship for this manuscript, had full access to all of the data in this study and
426 take complete responsibility for the integrity of the data and accuracy of the data analysis and
427 the work as a whole, and have given final approval to the version to be published.

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433 **Disclosures**

434 The authors declare they have no conflict of interest.

435 **Compliance with ethics guidelines**

436 Each of the 32 centers involved in the study obtained local ethics committee approval and
437 informed consent was obtained from all patients for being included in the study.

438 **Data availability**

439 The datasets generated during and/or analyzed during the current study are available from
440 the corresponding author on reasonable request.

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- 517

518 **TABLES**519 **Table 1.** Baseline patient characteristics. SD, standard deviation

Characteristic	N=1500
Mean ± SD age, years	64.8 ± 12.3
Gender, n (%)	
Male	810 (54)
Female	690 (46)
Mean ± SD Karnofsky Index score	61.1 ± 18.2
Place of assessment, n (%)	
Outpatient clinic	549 (37)
Day hospital	171 (11)
Home care	232 (15)
Hospice	47 (3)
Hospital inpatient ward	501 (33)
Primary tumor site, n (%)	
Lung	352 (22)
Urogenital	254 (17)
Gastrointestinal	276 (18)
Breast	201 (13)
Pancreas	129 (8)
Liver	16 (1)
Head and neck	97 (6)
Others	241 (15)
Disease, n (%)	
Loco-regional	250 (17)
Metastatic	1250 (83)
Previous anticancer treatment, n (%) ^a	1154 (79)
Care setting, n (%)	
Palliative care	289 (19)
Oncology	672 (45)
Pain therapy	526 (35)
Radiotherapy	13 (1)

All values are presented as mean ± SD or number of patients (proportion of patients)

a. data available in 1464 patients.

521 **Table 2** Patient characteristics associated with duration of breakthrough pain. BTP,
 522 breakthrough cancer pain; n, number of patients; SD, standard deviation.

Characteristic		n	Mean duration of BTP, minutes	SD	p-value
Disease	Locoregional	109	36.54	34.54	0.03
	Metastatic	395	44.64	41.34	
Primary tumor	Other	90	36.00	36.59	0.04
	Gastrointestinal/liver	89	42.36	38.99	
	Pancreas	56	55.09	46.35	
	Lung	99	42.26	40.41	
	Breast	68	38.18	33.75	
	Head and neck	16	50.38	58.09	
	Urogenital	86	45.74	39.70	
Anticancer treatment	No	97	37.24	32.97	0.05
	Yes	388	45.06	42.22	

523

524 **Table 3** Frequency of breakthrough pain according to care setting.

	Care setting					p-value
	Palliative care	Oncology	Radiotherapy	Pain therapy	All	
N	289	672	13	526	1500	
Type of BTP experienced, n (%)						
Mixed	113 (39)	411 (61)	6 (46)	364 (69)	894 (60)	
Neuropathic	8 (3)	63 (9)	0 (0)	15 (3)	86 (6)	<0.001
Nociceptive	168 (58)	198 (29)	7 (54)	147 (28)	520 (35)	

525

526 **Table 4.** Time to meaningful pain relief by treatment and other variables. BTP, breakthrough
 527 pain; FBST, fentanyl buccal sublingual tablet; FBT, fentanyl buccal tablet; FPNS, fentanyl
 528 pectin nasal spray; INFS, intranasal fentanyl spray; IV-M, intravenous morphine; OM, oral
 529 morphine; OTFC, oral transmucosal fentanyl citrate; SC-M, subcutaneous morphine; SD,
 530 standard deviation.

	Mean \pm SD time to pain relief, minutes	p-value
BTP treatment		
FBST	16.15 \pm 14.3	
FBT	13.78 \pm 11.0	
FPNS	10.99 \pm 8.6	0.012
INFS	10.64 \pm 5.2	0.012
IV-M	13.44 \pm 8.6	0.012
SC-M	15.36 \pm 10.2	
OM	18.84 \pm 12.1	
OTFC	12.97 \pm 5.4	
Other	27.73 \pm 18.1	
Place of assessment		
Outpatient clinic	23.08 \pm 18.0	
Day hospital	14.95 \pm 10.8	
Home	16.24 \pm 13.0	
Hospice	14.82 \pm 8.0	
Inpatient ward	14.05 \pm 11.0	<0.001
Primary tumor site		
Gastrointestinal-liver	15.29 \pm 11.7	
Pancreas	13.93 \pm 10.8	0.0193
Lung	16.11 \pm 16.0	
Breast	23.02 \pm 18.9	
Head and neck	14.33 \pm 10.7	0.0193
Urogenital	19.60 \pm 12.9	
Other	16.87 \pm 13.0	

532 **Table 5** Multivariate model for dissatisfaction. 95% CI, 95% confidence interval; BTP,
 533 breakthrough pain; FBST, fentanyl buccal sublingual tablet; FBT, fentanyl buccal tablet; FPNS,
 534 fentanyl pectin nasal spray; INFS, intranasal fentanyl spray; IV-M, intravenous morphine; OM,
 535 oral morphine; OR, odds ratio; OTFC, oral transmucosal fentanyl citrate; SC-M, subcutaneous
 536 morphine.

	OR (95% CI)	p-value
BTP treatment		0.0002
Other vs FPNS	1.98 (1.42 to 2.76)	
FBST vs FPNS	1.51 (1.03 to 2.21)	
FBT vs FPNS	1.31 (0.86 to 1.99)	
INFS vs FPNS	0.41 (0.14 to 1.24)	
IV-M vs FPNS	0.47 (0.22 to 1.00)	
SC-M vs FPNS	0.99 (0.50 to 1.94)	
OM vs FPNS	1.35 (0.93 to 1.95)	
OTFC vs FPNS	1.65 (0.95 to 2.88)	
Place of assessment		0.04
Day hospital vs outpatient clinic	0.71 (0.45 to 1.13)	
Home care vs outpatient clinic	0.29 (0.10 to 0.85)	
Hospice vs outpatient clinic	0.52 (0.15 to 1.72)	
Inpatient vs outpatient clinic	0.72 (0.51 to 1.03)	
Previous anticancer treatment vs no previous anticancer treatment	1.41 (1.06 to 1.87)	0.0166
Care setting		0.0011
Palliative care vs oncology	0.84 (0.29 to 2.44)	
Radiotherapy vs oncology	1.58 (0.54394 to 4.59)	
Pain therapy vs oncology	0.53 (0.37561 to 0.74)	

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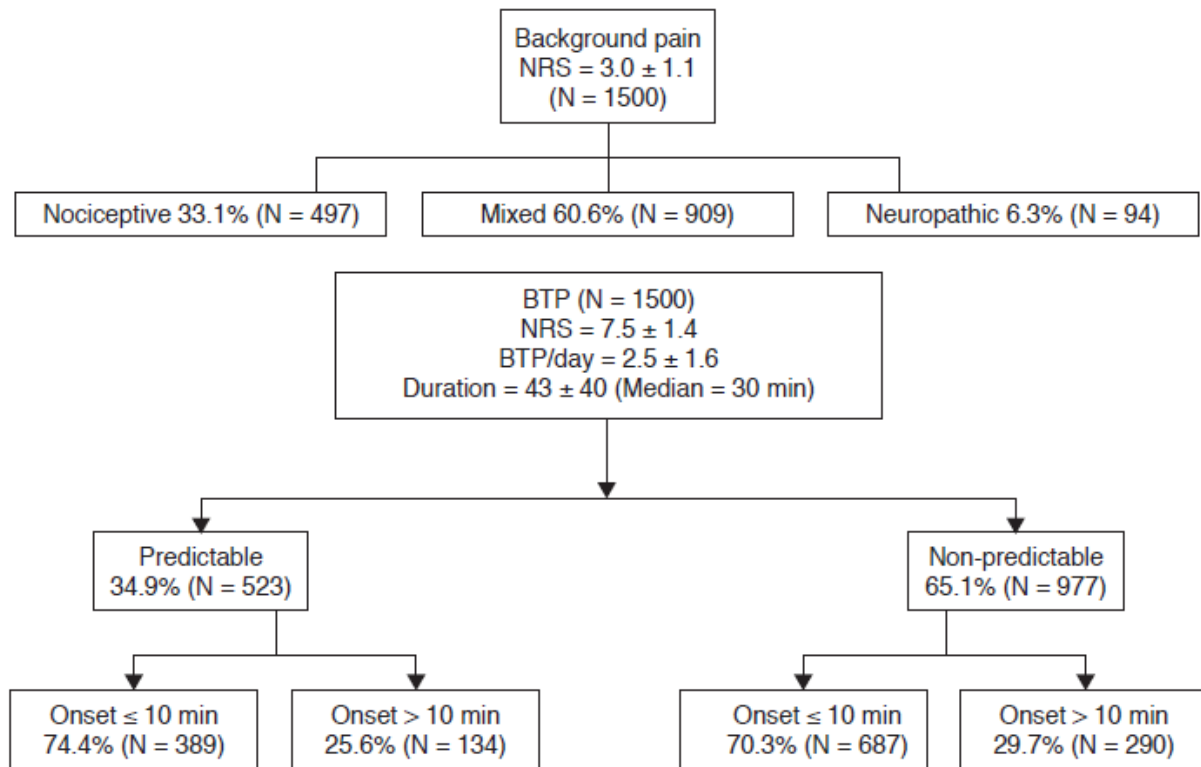
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544 Figure 1. Percentages of patients with baseline pain and characteristics of BTP

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546

BTP: breakthrough pain; NRS: numerical rating scale.