**Name of journal: World Journal of Gastroenterology**

**ESPS Manuscript NO: 6122**

**Columns: REVIEW**

**Celiac plexus neurolysis in pancreatic cancer: The endoscopic ultrasound approach**

Seicean A.EUS celiac neurolysis in pancreatic cancer

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**Received:** October 3, 2013  **Revised:** November 1, 2013

**Accepted:** November 18, 2013

**Published online:**

**Abstract**

The pain in pancreatic cancer is often the major problem of treatment. Administration of opioids is frequently limited by side effects or insufficient analgesia. Endoscopic ultrasound-guided celiac plexus neurolysis (EUS-CPN) represents an alternative for the palliative treatment of visceral pain in patients with pancreatic cancer. This review focuses on the indications, technique, outcomes of EUS-CPN and predictors of pain relief. EUS-CPN should be considered as the adjunct method to the standard pain management. It moderately reduces pain in pancreatic cancer, without eliminating it. Nearly all patients need to continue opioid use, often at a constant dose. The effect on the quality of life is controversial and survival is not influenced. The approach could be done in central position of the celiac axis, which is easy to perform, or in the bilateral position of the celiac axis, with similar results in terms of pain alleviation. The EUS-CPN with multiple intraganglia injection approach seems to have better results, although extended studies are still needed. Further trials are required to enable more confident conclusions on timing, quantity of alcohol injected and the method of choice. Severe complications have rarely been reported and great care should be taken in choosing the site of alcohol injection.

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**Key words:** Endoscopic ultrasound; Celiac neurolysis; Pancreas; Cancer; Pain

**Core tip:** Endoscopic ultrasound-guided celiac plexus neurolysis should be considered as the adjunct method to the standard pain management. It moderately reduces pain in pancreatic cancer, without eliminating it. Nearly all patients need to continue opioid use, often at a constant dose. Central technique is easy to perform, but intraganglia injection seems to give better results. This review focuses on methods of celiac neurolysis, with details about the endoscopic ultrasound guided celiac plexus neurolysis, indications, outcomes, which comprises efficacy, safety, novel techniques, and predictors of pain response.

Seicean A. Celiac plexus neurolysis in pancreatic cancer: The endoscopic ultrasound approach.

**Available from:**

**DOI:**

**INTRODUCTION**

The incidence of pancreatic cancer has increased over the last decade[1,2]. Few of the patients are diagnosed at an resectable stage (12%-20%)[3,4] and vascular resection during duodenopancreatectomy increases the 30-d postoperative morbidity and mortality rate[5]. For pancreatic cancer patients, the standardized net survival at 5 years is 6% for men and 10% for women[6]. Thus, palliative treatment is crucial in management. In this context, one of the most important symptoms to treat is pain. In the initial phase, the pain is visceral, but with disease progression, somatic pain may occur, especially due to the peripancreatic invasion of neural structures or muscles. Medicinal palliation of pain from pancreatic cancer begins with non-opioid drugs, such as paracetamol, stepping up to opioids, such as tramadol, and, eventually, more powerful opioids, such as morphine or fentanyl, as necessary. However, the dosage of opioid medication sometimes reaches a limit level due to side effects, such as nausea, constipation, somnolence, addiction, confusion or respiratory depression, and failure in achieving adequate analgesia. In these situations, neurodestructive methods involving the main pancreatic pain pathways, such as celiac block or thoracoscopic splanchnicectomy, seem efficient.

The celiac ‘plexus’ is the largest plexus of the sympathetic nervous system, innervating the upper abdominal organs (pancreas, diaphragm, liver, spleen, adrenal glands, kidneys, abdominal aorta, mesentery, stomach, small bowel, ascending colon and the proximal portion of the transverse colon). The celiac plexus is situated within the retroperitoneal space posterior to the stomach and pancreas, close to the celiac axis, and it is separated from the vertebral column by the crush of the diaphragm. It comprises a dense network of ganglia around the aorta, with considerable variability in size (0.5-4.5 cm), number[7-11] and position (from the T12-L1 disc space to the middle of the L2 vertebral body). The left celiac plexus is typically located more caudally than its counterpart on the right. Celiac neurolysis may target either the plexus or the ganglia.

The preganglionic sympathetic fibres of the celiac plexus are grouped into the greater (T5-10), lesser (T10-11) and the least (T12) splanchnic nerves, and the plexus also receives parasympathetic fibres from the celiac branch of the right vagus nerve. All of these fibres are interrupted during thoracoscopic splanchnicectomy performed under general anaesthesia (Figure 1). This review focuses on the following aspects: methods of celiac plexus neurolysis, with details about the endoscopic ultrasound guided celiac plexus neurolysis (EUS-CPN), indications, outcomes, which comprises efficacy, predictors of pain response, safety and novel techniques.

**CELIAC PLEXUS NEUROLYSIS**

This involves chemical destruction of celiac ganglia and corresponding neural pathways by injecting dehydrated alcohol into the network of the celiac plexus. The result is moderate neuronal degeneration associated with residual fibrosis[7].

The initial method involved a posterior approach, accomplished under guidance by fluoroscopy or computed tomography (CT). The pain level at 12 wk after the procedure was significantly lower than with systemic analgesic therapy [8]. Unfortunately, in some cases the pleura or neural structures were accidentally touched by the needle, with subsequent development of serious side effects such as pneumothorax and paraplegia, respectively[8].

As a consequence, the anterior approach came to be considered a better option for CPN, accomplished either transcutaneously-under ultrasound (US) guidance (developed in 1995[9]), CT guidance or endoscopic US (EUS) guidance-[introduced in 1996[10]] or, more invasively, by means of surgery. More recently, the laparoscopic technique has been implemented[11,12].

The advantages of the EUS approach are the fine orientation of the needle above or lateral to the celiac trunk and the real-time performance of the procedure, under Doppler control of vessel interposition. In addition, the technique is easy, requiring only 2-3 min immediately after the staging or sampling of an inoperable pancreatic tumour. Better results can be expected owing to the better orientation of the needle, compared to the US or CT approach, and the real-time accomplishment of the procedure.

The technique consists in preprocedural hydration with 500 mL saline, followed by CPN performed with the patient in left lateral position, under either general anaesthesia with propofol, or deep intravenous sedation with 2-4 mg of midazolam. Some endosonographers favour antibiotic prophylaxis, avoiding a retroperitoneal abscess[13-16], although alcohol is considered to be a bactericidal agent[17]. Bacterial translocation from gut might be reduced by performing a single needle pass and by avoiding simultaneous gastric acid suppression treatment[17,18]. After colour Doppler assessment of vessel-gut interposition, a therapeutic linear-array echo-endoscope is used and the puncture site is chosen. Proximity to the diaphragm should be avoided, because of the potential for immediate pain, due to the spread of alcohol. The devices used are 22-G or 19-G needles, or preferably fenestrated 20-G needles especially designed for EUS-CPN [Cook Medical, Winston-Salem, NC, United States)]. Recently, the use of a forward-viewing echo-endoscope has been reported in five patients[19].

For central injection, which is easier to perform, the needle is advanced above the celiac trunk, in the space between the aorta and the origin of the celiac axis. If bilateral injection is chosen, the echo-endoscope, situated above the celiac axis, is rotated to one side until the origin of the celiac axis is no longer seen, and half of the entire solution is injected; the procedure is then repeated on the opposite side.

When ganglia are targeted, the echoendoscope is rotated clockwise and celiac ganglia are found above the celiac trunk, alongside the trunk, and below the trunk, just above the superior mesenteric artery takeoff[20]. The ganglia are small hypoechoic nodules with hyperechoic foci in the center. Sometimes their interconnection can be seen. In large ganglia, thin linear hypoechoic lines arising from the edges of the ganglion are suggestive of small neural fibers[20]. The rate of ganglia detection varies between 79%-89% and it may also vary among endosonographers (65%-97%)[21-23]. As many ganglia as possible should be injected. The actual recommendations are to start the injection in the central part of the ganglia for those within 1 cm in diameter, or in the deepest part of the larger ganglia, and to perform the injection during the withdrawal of the needle , but only inside the ganglia[24].

Following the Doppler assessment of the area, aspiration is performed in order to rule out placement of the needle inside a vessel, which may lead to severe complications (Figure 2). Any lack of resistance during injection might suggest that the vascular space has been punctured, the needle should be withdrawn, and the aspiration test repeated. The injection starts with 3-10 mL of a local analgesic to prevent transient pain exacerbation induced by the neurolytic agent. Lidocaine 1-2%[13,14,20,25], or a better analgesic, such as bupivacaine 0.25%-0.75%[10,23,26-28] can be used..Subsequently,10-20 mL of a neurolytic agent (98% dehydrated alcohol) is injected and a hyperechoic cloud is seen in the area of the needle tip, as the substance spreads.

All patients must be kept under close observation for 2 h after the procedure, to monitor blood pressure, heart rate and temperature, and to identify any immediate complications.

Technical difficulties may occur in some cases because the anatomical landmarks could not have been be properly visualized. For example, during bilateral technique, after one side injection, the alcohol spreads and impedes the view of the opposite side. Sometimes, the celiac plexus region cannot be reached with the needle, as in patients with cachexia who have very little fat tissue around the aorta, or when the diaphragm insertion is too close to the celiac trunk.

Other approaches such as thoracoscopic splanchnicectomy, EUS-guided direct celiac ganglion irradiation with 125I seeds or radiofrequency ablation could be considered the alternative methods for celiac destruction[11,29,30].

**INDICATIONS OF EUS-CPN**

The NCCN guidelines, version 2.2012 for pancreatic adenocarcinoma, recommend EUS-CPN for the treatment of severe tumour-associated pain. This is useful especially when intolerable adverse efects of opioid therapy occur, such as drowsiness, delirium, dry mouth, anorexia, constipation, nausea and vomiting, or an analgesic “ceiling” is seen due to neurotoxicity. In the case of jaundice caused by an unresectable pancreatic head tumour, biliary drainage should be offered first, followed by EUS-CPN if pain persists[31].

Relative contraindications to EUS-CPN include difficult access, due to anatomical distortion from previous surgery or congenital malformations. The absolute contraindications for EUS-CPN are the same as for any other invasive procedure: coagulopathy, platelets < 50000, and patients who are unable or unwilling to cooperate[32].

**OUTCOMES**

***Efficacy for EUS-CPN***

The main goals when performing EUS-CPN are the pain alleviation and the improvement of the quality of life. This procedure added no benefits regarding survival in two randomized controlled trials [13,27].

Although the quality of life was unchanged after CPN in one randomized trial [16], there are reports of improvement of parameters of the quality of life such as functional status, work capability, sleep, and enjoyment of leisure activities[14,26]. The occurrence and duration of terminal delirium have also been reported as reduced after this procedure[33].

The assessment of pain intensity in chronic cancer patients uses different measurement scales. Visual analogue scales proved to be less suitable in old patients with opioids use, due to limited communication skills and cognitive impairment during the last days of life, making self-reporting of pain more difficult [34]. Numerical rating scale is preferable in assessing cancer pain exacerbations to verbal rating scales [35].As a result, observation of pain-related behaviours and discomfort is indicated in patients with cognitive impairment, in order to assess the presence of pain[36], and multidimensional questionnaires which evaluate the pain intensity together with other parameters of interference with pain are useful[26].

Although the real benefit of EUS-CPN compared to placebo has not been studied, pain relief after the procedure varies between 45%-94% in different papers (Table 1). Two subsequent meta-analyses showed a mean rate of pain alleviation of 72%-80%, with a much lower rate of complete pain response[24,42,43]. However, many of the patients still require the same dose of analgesic and EUS-CPN should be considered as an adjunct method to the standard pain management{24]. The post-neurolytic residual pain could be related to the non-visceral pain, due to the invasion of the muscles or surrounding connective tissue, but factors concerning the technique used (type of technique, quantity of alcohol injected, timing of the procedure) were extensively studied.

The type of technique used for obtaining the best response is still controversial. Eleven studies on the central or bilateral technique have been published to date, showing a pain alleviation rate of 50%-88% at 1-14 wk after the procedure (Table 1).Bilateral technique, used in six of these studies, was associated with a rate of pain alleviation of 45%-88%, while central technique showed 68%-72% alleviation. To date, only one randomized controlled trial has compared the central and bilateral techniques of EUS-CPN and showed no difference in duration of pain relief (11 vs. 14 weeks), complete pain relief (2/29 *vs* 2/21 patients) or reduction in pain medication (9/29 *vs* 7/21 patients)[13]. The choice between the central or bilateral technique remains difficult, depending on the personal skills and the experience of every endosonographer. Our experience has showed good results with central technique, which we consider easier to perform[26].

EUS-guided direct ganglia neurolysis, first reported by Levy in 2007, showed much better results in terms of pain alleviation (7 of 17 patients, 94%) at 2-4 wk; the known side effects – diarrhoea or hypotension – were noted. For the first time, long-lasting postprocedural pain (2.2 d) was reported in 7 of 17 patients [15]. One-week follow-up of pain alleviation showed again better results for EUS-CPN at the celiac ganglia compared to EUS-CPN at the celiac trunk region using bilateral injection (67.5% *vs* 33%)[14]. However, this technique was used in only few studies. One randomized control trial compared direct ganglia neurolysis with central neurolysis. The positive response rate at day 7 and the complete response rate was higher in the ganglia neurolysis group (75.5%*vs* 45.5%, respectively 50% *vs* 18.2%)[23]

***Early vs late injection***

A randomized double-blind controlled trial in 96 patients showed that CPN was effective in pain reduction at 1-month and 3-month follow-up, but opioid consumption was constant -- although it increased in the control group[27]. In the group of patients without radiochemotherapy, pain was significantly reduced and the need for increased opioids was prevented. In patients with radiochemotherapy, on the other hand, pain was significantly reduced only at 3 months of follow-up. The authors concluded that this technique would be effective only for patients who refuse, or are ineligible for radiochemotherapy[27].

***Amount of ethanol injection***

The majority of cases were done using 10-20 mL alcohol[13-15,26,27]. Only one study compared the results when 10 or 20 mL alcohol injection was used during intraganglia or central injection and no difference in pain alleviation was noted[14].

***Repetitiveness of the procedure***

The benefit of repeated EUS-CPN was studied in 24 patients and results are less encouraging. The rate of successful pain relief was much lower than for the first EUS-CPN (29% *vs* 67% at 1 mo follow-up), and disease progression was a factor which limited the response[44].

**EFFICACY FOR PERCUTANEOUS CPN AND ALTERNATIVE APPROACHES**

Two important meta-analyses of the percutaneous approach have been published. The first one included 1117 patients, 63% of them with pancreatic cancer, in whom bilateral X-ray-, US- or CT-guided neurolysis was performed. Pain alleviation at 2 weeks was excellent, 90% relief was recorded at 3 months after the procedure, and 70%-90% of patients experienced pain relief right up to the time of death. Transient pain was seen in most of the patients under study (96% in two of the studies analysed), transient diarrhoea in 44%, and transient hypotension in 38%. Severe neurological side effects were noted in 5 of the 268 patients (1%)[45]. A second meta-analysis, including 358 patients with CT-guided CPN, from six randomized controlled trials, showed a limited advantage in pain alleviation at 4 and 8 weeks (0.42 and 0.44 respectively on a visual analogue scale of 0-10), but opioid consumption was significantly lower, with fewer side effects[46].

To date, only one published randomized trial has compared the efficacy of CPN and thoracoscopic splanchnicectomy, and the results were not significant compared to the control medical management group. The main limitation of the study was the small number of patients included in each arm of the study[11].

EUS-guided direct celiac ganglion irradiation with 125I seeds was performed in 23 patients, with significant pain reduction 2 weeks later. Initial pain exacerbation was seen in 26% of the patients, but no major complications occurred up to the time of death [29].

Radiofrequency ablations of pancreatic mass and celiac plexus have been reported as successful in the treatment of chronic pain [30].

***Predictors of response rate***

A retrospective study compared the results of ganglia injection with those of non-direct ganglia injection (40 *vs* 24 patients). The median number of visualized ganglia was two. The pain response rate was 50% at 1 wk, 77% at 30 d, and opioid consumption was 57% lower at 1-week follow-up. Pain alleviation was significantly lower for patients in whom the ganglia were not visualized, and it was also lower, albeit not significantly so, for tumours located in the body or tail of the pancreas, for large tumours and for patients with severe pain at presentation[20].

A second study on 47 patients with central-injection CPN showed 68% pain alleviation at 1 week. The predictors of poor pain alleviation were direct invasion of celiac ganglia and left diffusion of the neurolytic agent[37].

***Safety***

Many complications have been described for EUS celiac block indicated for chronic pancreatitis, and some of these complications have been seen in EUS-CPN for pancreatic cancer, such as transient diarrhoea (4%-15%) and transient hypotension (1%)[14,18,47,48] or alcohol intolerance. Nowadays, it is considered that the potential immediate complications are rare, such as hypotension, tachycardia, initial pain enhancement, severe bleeding and paraplegia. The late side effects include diarrhoea, hypotension, fever and paraplegia[48].

Recently, severe complications of EUS-CPN have been reported in individual cases, but they have to draw the endosonographers’ attention (Table 2). Permanent lower paraplegia, due to spinal cord infarction, was noted in one patient; the mechanism was considered to be alcohol diffusion via the left T12 intercostal artery towards the anterior spinal artery, or vasospasm caused by alcohol or acute thrombosis due to injection of a high volume into the celiac area[50,51]. Injury of the lumbar artery leading into the artery of Adamkiewicz could be involved. This major artery originates from the aorta, varies in position from T7 to L4, supplies the lower two-thirds of the anterior spinal artery, and it is anatomically closely related to the celiac ganglion[51]. The clinical manifestations, reported 14 h after the procedure, comprised motor weakness, decreased pain and temperature sensation below T7-L1, and detrusor atony. Prolonged periprocedural hypotension may have played a part[51]. Extreme caution should be taken concerning the placement of the needle tip, including Doppler US examination of the area and aspiration before injection.

Another complication was thrombosis of the celiac trunk, with wall thickening and bubble-like pneumatosis of the stomach, duodenum, jejunum, ileal loops and ascending colon. Signs of hepatic infarction of segment I and III, near-total right kidney, as well as splenic infarction were discovered, and the evolution was fatal. The explanation was the sclerosing effect of alcohol after injection[49]. Alcohol neurolysis for treatment of chronic pancreatitis has been recorded as leading to necrosis and perforation of the stomach and aorta with lethal outcome[52], as well as splenic, gastric and pancreatic infarction[54]. The infarction of the liver, spleen, stomach, and proximal small bowel  after celiac neurolysis for pancreatic metastasis was reported in one case, as vasospasm resulted from the diffusion of ethanol into the celiac artery[53].

Retroperitoneal abscess have been previously noted in chronic pancreatitis patients with triamcinolon injection, especially after gastric acid supression therapy[17,18], but one case has been described in pancreatic cancer, too[16]. However, the antibiotic prophylaxis is not done in many studies[23,25-28].

Initial pain exacerbation after EUS-CPN was reported in up to 29%-34% of cases[15,22]. Previous studies considered initial pain exacerbation as a sign of greater pain relief at follow-up[15], but this was not confirmed in further studies[22].

***Novel techniques***

In the attempt to improve the technique, Sakamoto *et al*[40] used broad plexus neurolysis near the superior mesenteric artery with the aim of administering the neurolytic agent to a larger number of ganglia. The authors checked the spread of the neurolytic agent around the celiac axis and showed that the new technique achieved neurolysis in five or six areas in a higher proportion of patients than the previous method. With regard to the pain levels at 7 d and 30 d, significant reduction was obtained for five and six areas of neurolytic agent diffusion, but not for three or four areas. However, this study had some limitations: there were methodological problems, the physicians’ experience increased during the study, not all patients (only 60 of 67) had pancreatic cancer, and the overall success rate for pain alleviation was only 50% at 30 d, the lowest rate ever reported at that time[38].

**CONCLUSION**

EUS-CPN should be considered as the adjunct method to the standard pain management. It moderately eases pain in pancreatic cancer without eliminating it completely. Nearly all patients need to continue opioid use, often at a constant dose. Multicentre randomized controlled trials are required to provide more reliable conclusions on timing, quantity of alcohol injected and the method of choice. Until then, in the light of the severe complications reported recently, great care should be taken into consideration when choosing the site of alcohol injection.

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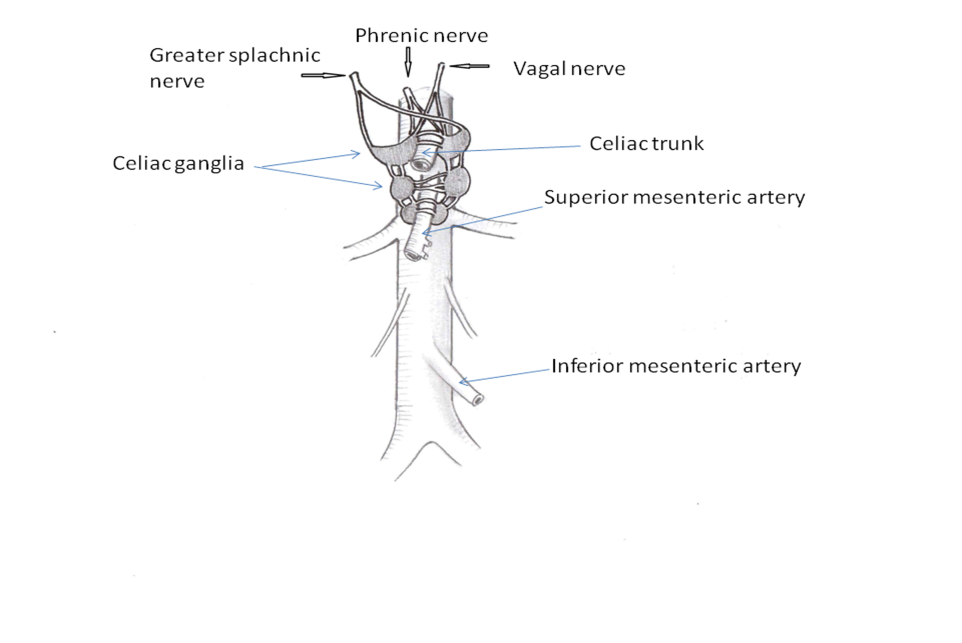
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**P-Reviewers:** Katanuma A, Iglesias-Garcia J, Meister T **S-Editor:** Qi Y

**L-Editor: E-Editor:**

**Figure 1 Anatomy of the celiac area (courtesy of Dr. C. Gombosiu).**



**Figure 2 Endoscopic ultrasound images showing the position of the needle above the celiac.**

**Table 1 Pain relief and techniques used in patients with endoscopic ultrasound-guided celiac plexus neurolysis**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Ref** | **No of patients** | **Pain evaluation** | **Technique of EUS-CPN** | **Follow-up period(wks)** | **Pain alleviation** |
| Doi *et al*[23] 2013 | 68 | Numeric rating scale | Ganglia *vs* central | 1 | 73% *vs* 45% |
| LeBlanc *et al*[14] 2013 | 20 | Numeric rating scale | Ganglia + central | 6 | 90% |
| Seicean *et al*[26] 2013 | 32 | Brief pain inventory | Central | 2 | 75% |
| Wiechowska-Kozłowska *et al*[25] 2012 | 29 | Numeric rating scale | Central + bilateral | 8-12 | 76% |
| Wyse *et al*[27] 2011 | 48 | Likert scale | Bilateral | 12 | 60.7% |
| LeBlanc *et al*[13] 2011 | 50 | Numeric rating scale | Centralvs bilateral | 14 | 69 *vs* 81% |
| Iwata *et al*[37] 2011 | 47 | Visual analogue scale | Central | 1 | 68.1% |
| Ascunce *et al*[20] 2011 (20) | 64 | Numeric rating scale | Ganglia or bilateral | 1 | 50% |
| Sakamoto *et al*[38] 2010 | 67 | Visual analogue scale | Under celiac trunk | 4 | 33%-93% |
| Sahai *et al*[39] 2009 | 160 | Visual analogue scale | Central versus bilateral | 1 | 70% *vs* 45% |
| Ramirez-Luna *et al*[40] 2008 | 10 | Visual analogue scale | Central | 4 | 72.2% |
| Levy *et al*[15] 2008 | 17 | General descriptors | ganglia | 4 | 94% |
| Tran *et al*[28] 2006 | 10 | Numeric rating scale | Central | Not stated | 70% |
| Gunaratnam *et al*[41] 2001 | 58 | Visual analogue scale | Bilateral | 24 | 78% |
| Wiersema *et al*[10] 1996 | 30 | Visual analogue scale | Bilateral | 12 | 79%-88% |

EUS-CPN: Endoscopic ultrasound-guided celiac plexus neurolysis.

**Table 2 Immediate and late complications reported for endoscopic ultrasound-guided celiac plexus neurolysis in adenocarcinoma patients**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Ref.** | **No. of procedures** | **Complications** | **Indication** | **Technique** | **Substance** |
| Muscatiello *et al*[16] | 1 | Retroperitoneal abscess | PC | Not stated | Alcohol + bupivacaine |
| Gimeno-Garcia *et al*[49] | 1 | Celiac axis infarction, kidney, splenic, hepatic infarction, death | PC | Bilateral | Alcohol + bupivacaine |
| Fujii *et al*[50] | 1 | Anterior spinal cord infarction with lower paraplegia | PC | Ganglia + central | Alcohol + bupivacaine |
| Wiechowska-Kozłowska *et al*[25] | 29 | Hypotension-1  Pain exacerbation-2  Transient diarrhoea-3 | PC | Bilateral + central | Alcohol + bupivacaine |
| Mittal *et al*[52] | 1 | Anterior spinal cord infarct with lower paraplegia | PC | Ganglia + central | Alcohol + bupivacaine |
| O’Toole *et al*[17] | 31 | Hypotension-1 | PC | Bilateral | Alcohol + bupivacaine |
| Levy *et al*[15] | 17 | Pain exacerbation-2 | PC | Ganglia | Alcohol + bupivacaine |
| Leblanc *et al*[14] | 20 | Lightheadedness-1  Transient diarrhea-2  Transient nausea and vomiting-3 | PC | Central+  ganglia | Alcohol+ bupivacaine |
| Doi *et al*[23] | 68 | Transient hypotension-3  Upper gastrointestinal bleeding-1  Pain exacerbation-17  Transient diarrhea-5  Inebration-2 | PC | Central+  ganglia | Alcohol+  bupivacaine |
| Jang *et al*[53] | 1 | Liver and splenic infarction, ischemia of the stomach and small bowel | Pancreatic metastasis | Central | Alcohol+bupivacaine+triamcinolon |

PC: Plexus neurolysis in adenocarcinoma.