Video Capsule Endoscopy in Patients with Chronic Abdominal Pain with or without Associated Symptoms: A Retrospective Study

Jeremy Egnatios¹, Khushboo Kaushal¹,², Denise Kalmaz¹,²,³, Amir Zarrinpar¹,²,³*

¹ School of Medicine, University of California San Diego, La Jolla, California, United States of America,
² Department of Medicine, University of California San Diego, La Jolla, California, United States of America,
³ Division of Gastroenterology, University of California, San Diego, La Jolla, California, United States of America

* azarrinpar@ucsd.edu

Abstract

Background
Chronic abdominal pain (CAP) is a common indication for gastroenterology referrals. More insidious causes of CAP isolated to the small bowel, such as malignancies and Crohn’s disease, are rising in incidence and causing more gastroenterologists to evaluate their patients with video capsule endoscopy (VCE). However, the role of VCE in patients with CAP is still unclear.

Aims
We assessed the efficacy of VCE in patients with CAP and whether it led to findings that contributed to disease management and meaningful interventions.

Methods
This retrospective study evaluated 607 capsule endoscopy studies at an open referral endoscopy unit. Ninety of the studies were for CAP. These studies were compared to those performed for other indications to compare diagnostic yield. In addition, we investigated whether VCE led to an intervention that improved clinical outcomes.

Results
Overall, the number of abnormal findings in CAP patients was significantly lower than VCE performed for other indications (24.4% vs 39.0%, respectively p = 0.009). When patients with CAP presented with other pertinent clinical findings (e.g. nausea, weight loss, anemia, history of inflammmatory bowel disease, etc.), the likelihood of an abnormal finding increased to a level that was not different from those who received VCE for other indications (27.1%, p = 0.10). The findings from VCE lead to changed management and improved outcomes in 16.2% of CAP patients with associated symptoms. However, the subgroup that benefited the most were those who had a prior history of Crohn’s disease. Patients with
CAP who did not have any associated symptoms continued to have a significantly lower abnormal finding rate compared to those who received VCE for other indications (19.4%, p = 0.03) and VCE rarely led to a change in management that would improve outcomes (5.6%).

Conclusions

VCE for CAP has a lower rate of abnormal findings than other indications. However, VCE is a useful diagnostic tool that can help provide a possible etiology of CAP in patients with associated symptoms. However, a change in management from VCE is likely to be limited to those with a history of Crohn’s disease.

Introduction

Chronic abdominal pain (CAP) is a common gastrointestinal complaint, with a prevalence of up to 21% in the general population [1-3]. It is a significant economic burden, with up to $20 billion in direct and indirect costs [4]. CAP is the 8th most common complaint of all patients presenting in primary care for a health issue and one of the most common reasons for gastroenterology clinic visits [5, 6]. There are many etiologies for abdominal pain ranging from benign and functional disorders to malignancy and inflammatory diseases (IBD), making evaluation of these patients at times difficult. To date, there are no consensus recommendations or practice guidelines for the evaluation of CAP in adults.

Although esophagogastroduodenoscopy (EGD) and colonoscopy are commonly used to evaluate patients with CAP, the role of video capsule endoscopy (VCE) remains unclear. Studies evaluating the diagnostic yield of VCE in CAP patients are limited and inconsistent, with a great range in diagnostic yield (4–44%) [7, 8] and were almost exclusively done in tertiary care referral centers [9]. The studies rarely report whether VCE changed management or improved outcomes. Few studies report how the CAP population are different, both in demographics and their diagnostic yield, from those who received VCE for other reasons. Given the wide range in diagnostic yield, relative safety, and coverage by insurance providers, VCE remains commonly used in practice for the evaluation of patients with CAP [9].

A recent systematic review of studies evaluating the diagnostic potential of VCE in CAP concluded that its use is unclear and conflicting [9]. It also concluded that the current literature regarding the usefulness of VCE in CAP is incomplete considering differences in study parameters, inclusion/exclusion criteria, and varying definitions of outcome reporting. Other factors including different referral base, sample size, and patient characteristics vary greatly between studies. Hence, they limit the interpretation of studies as a whole and obscure the usefulness of performing VCE to evaluate patients with CAP. Finally, the most important aspect of using VCE in practice, whether a diagnosis affected management which improved outcome, is not reported in any previous study.

In this study, we present a retrospective study from the largest cohort of American patients. We investigated the use of VCE in patients with CAP referred to an open endoscopy center. These referrals included those by community physicians as well as from a tertiary referral center. The performance of VCE for CAP was compared with those who received VCE for other indications and the outcomes after diagnosis were also measured. We aimed to determine the yield of performing VCE in CAP and whether it significantly altered management.
Methods

Participants

The University of California, San Diego (UCSD) Human Research Protections Program Institutional Review Board approved this retrospective study. Since this was a retrospective study where patient-identifying information was not part of final analysis, no consent was required. UCSD Medical Center Advanced Endoscopy Unit is an open referral center, drawing patients from throughout San Diego and Imperial Counties. Between June of 2004 and July of 2013, 607 capsule endoscopy studies were performed. Studies were examined for primary and ancillary indications, stated by physician, as reason for referral. Patients were excluded if study evaluator determined it incomplete because of poor bowel prep, or if the capsule did not reach the cecum. The endoscopy referral center was not responsible for the workup of CAP and the referral center did not exclude patients who may have had incomplete diagnostic procedures prior to capsule studies. However, almost all (>99%) of the patients had received colonoscopy and EGD studies prior to VCE.

CAP was defined as any type of abdominal pain lasting greater than three months from onset. Patients with CAP were initially evaluated as a single group. With further analysis, they were allocated by their associated symptoms or relevant past medical history (Fig 1). Patients

![Flowchart with classification of patients with chronic abdominal pain (CAP), along with subgroup classification based on clinical presentation. Abbreviations: PMH (past medical history), CD (Crohn's disease), UC (ulcerative colitis).](doi:10.1371/journal.pone.0126509.g001)
were divided into two groups: CAP only (CAP-O; when no other symptoms or relevant PMH was listed), CAP with associated findings (CAP-A; patients had accompanying chronic nausea, vomiting, diarrhea, weight loss, gastrointestinal bleeding or past medical history of Crohn’s disease, ulcerative colitis, or malignancy). In the CAP-A group, thirteen patients had a past medical history of Crohn’s. We analyzed whether the patients with Crohn’s disease affected group characteristics and later analysis. All analysis was compared with and without the Crohn’s disease patients, with any significant change in findings noted in the results. All remaining non-CAP indications were grouped for control comparison.

Retrospective Review

All patient information was taken, retrospectively, from the UCSD electronic medical records. When available, patient data collected included: age, sex, body mass index (BMI), past medical history, relevant associated symptoms (such as diarrhea, weight loss, vomiting, etc.), current medications, indication for VCE studies, gastric and small bowel transit times, inpatient status, diagnostic findings, and patient outcome from diagnostic findings. All VCE studies were read by at least two independent gastroenterologists, including one who is highly experienced in small bowel disease and evaluation of small bowel pathology (DK). The gastroenterologists who read the studies were not blinded to the reason for referral. Studies were considered to be positive for an abnormal finding if any imaging deemed abnormal by endoscopist was in the report, regardless of whether the observed findings could be related to patient’s presenting symptoms.

Additional chart review was performed on patients with abnormal finding. Differences in demographics, referral source, and serum inflammatory markers were assessed. Their records were then evaluated for follow up and management for at least 6 months after VCE. We assessed whether successful interventions or disease management were a result of abnormal findings from VCE. Patient outcomes were determined by reviewing physician’s notes in the patient’s chart.

Sensitivity Analysis

In order to investigate whether the benefit from interventions for VCE was comparable to that observed for other indications of VCE (i.e. GI bleed), we performed a sensitivity analysis. Patients who received a VCE for obscure gastrointestinal bleeding (VCE-GIB patients) were used as a separate cohort for this analysis. This subset of patients was then categorized by age and gender with no other information available to the investigator. Each patient with CAP was then matched to VCE-GIB patients in a 1:2 ratio (that is 2 VCE-GIB patients for each 1 CAP patient) for all patients that had follow up. Hence we made two separate cohorts, VCE-GIB-mA, who were matched to the CAP-A patients, and VCE-GIB-mO, who were matched to the CAP-O patients. As before, chart review for at least six months following VCE study was performed on the VCE-GIB patients and any information on intervention, further study, outcome, or resolution was recorded.

Statistical Analysis

Statistical analysis was performed using the Statistical Package for Social Science (SPSS) version 20.0 (SPSS, Inc., Chicago, IL) and GraphPad Prism 5 (GraphPad Software, La Jolla, CA). Student t-test was used to compare continuous variables (e.g. age, BMI, etc.). Chi-square analysis was used to compare categorical variables (e.g. abnormal findings, interventions, gender). Given the limited availability of VCE specialists and the referrals seen in this study, intent to treat analysis was applied to all comparisons, as we felt this best evaluated the true clinical impact of VCE. For all of statistical tests, an alpha level of less than 0.05 (two-tailed) was set for significance.
Results

Demographics

During the nine year period of the study, 607 capsule studies were performed, with 510 meeting the inclusion criteria. The majority of excluded studies, 85 out of 97 (88%), were due to the capsule not reaching the cecum. Ninety studies were for CAP, with 31 performed in patients who did not have any other symptoms. The remaining 59 studies were performed in patients who complained of CAP along with other associated symptoms (Fig 1). Patients who received VCE for other indications (e.g. iron deficiency anemia, overt bleeding, polyposis syndrome, etc.) were grouped into one cohort as a control for comparison (non-CAP; S1 Fig). There were demographic differences between patients with CAP and those in our control, non-CAP cohort (Table 1). Patients with CAP were significantly younger (43.7 years vs 55.5 years, \( p = 3.0 \times 10^{-5} \)) and more likely to be women, (66.7% v. 51.0%, \( p = 0.001 \)). None of the patients in the CAP group had a retained capsule or other complication from VCE.

Abnormal Finding Rate

VCE studies performed for CAP were less likely to yield an abnormal finding compared to the control group, (24.4% vs 39.0%, respectively, \( p = 0.009 \)). Since the patients with CAP are a heterogeneous group, we subdivided them into two groups: (1) CAP only (CAP-O, 31 patients) and (2) CAP with associated symptoms (CAP-A, 59 patients) (see methods for description of each group; Fig 1). On additional investigation of clinical presentation of CAP, no significant differences were found in rates of abnormal findings between the two CAP subgroups (19.4% for CAP-O; 27.1% for CAP-A; Fig 2). However, only the CAP-O group had significantly lower abnormal yield than the control group (\( p = 0.02 \)).

Since other symptoms appeared to affect the rate of abnormal findings, we investigated whether more objective measures, such as serum biomarkers, would as well. Specifically, inflammatory markers, such as CRP, ESR, and leukocytosis, as well as rheumatological markers such as ANA were analyzed for patients with a positive abnormal finding on VCE and those without any abnormal findings on VCE. However, none of these serum makers were predictive of an abnormal finding in CAP patients (S1 Table).

Community versus Tertiary Care Referral

Because our retrospective data was collected from an open endoscopy center, we compared patients referred from community physicians (47 patients; 52.2% of patients with CAP) and

| CAP-O (n = 31) | CAP-A (n = 59) | Total (n = 90) | Total (n = 420) | p-value* |
|----------------|----------------|----------------|----------------|---------|
| Age (yr)       | 43.1 (±2.3)    | 44.6 (±2.7)    | 43.7 (±1.7)    | 55.5 (±0.93) | 0.00003a |
| Gender (%)     | 58.1%          | 71.2%          | 66.7%          | 51.0%    | 0.001b   |
| BMI (kg/m²)    | 24.0 (±0.8)    | 24.5 (±0.5)    | 24.1 (±1.0)    | 26.5 (±0.5) | 0.03a   |
| % Inpatient    | 3.1%           | 19%            | 13.2%          | 19.1%    | 0.2b     |
| % Abnormal Finding | 19.4%      | 27.1%          | 24.4%          | 39.0%    | 0.009b   |

CAP-O represents CAP with no other symptoms. CAP-A represents CAP and any associated symptoms (See Methods for description).

*P value represents analysis between all patients with CAP and Non CAP groups.

a Student t test.
b \( \chi^2 \) test.

doi:10.1371/journal.pone.0126509.t001

Video Capsule Endoscopy in Chronic Abdominal Pain

PLOS ONE | DOI:10.1371/journal.pone.0126509 April 20, 2015 5/16
those from the tertiary care facility (43 patients; 47.8% of patients with CAP). Community physicians referred the majority of patients with CAP and no other symptoms, (20 patients; 64.5% of CAP-O). However, patients referred from community physicians were much more likely to have abnormal findings on VCE studies compared to those from tertiary care facility (34.0% v 14.0%, respectively, p = 0.003) (Fig 3A).

### Additional Testing After VCE

We also investigated the use of other diagnostic procedures for patients with CAP. Interestingly, we found that CAP-O patients were more likely to have had gastric emptying studies performed compared to the control cohort (16.5% v 4.8%, respectively, p = 4.0x10^-5). However, they were not likely to have abnormal findings from those studies as well, questioning need for such an extensive work-up (Fig 3B). The CAP-A group did not have a significant difference in this particular measure when compared to the control cohort.

### Post-VCE Management and Intervention

In order to further understand the role of VCE in the evaluation of patients with CAP, we investigated whether VCE affected management and interventions. We had follow up data for 61 (67.8%) of patients with CAP, including 18 (58.1%) of CAP-O group and 43 (72.9%) of CAP-A group. Follow up data was available for the vast majority of patients with abnormal findings (16/22; 72.7%) for at least 6 months after VCE. In patients with CAP, 24 out of the 61 (39.3%) had improvement of symptoms on follow-up. The majority, 13 patients (54.2%), were due to spontaneous resolution. Of the CAP group, 11 patients (47.8%) improved from an intervention after VCE. Of the 24 patients whose symptoms resolved after VCE, 19 (79.2%) were from the CAP-A group, 10 (52.6%) of whom benefited from a post-VCE intervention (Table 2, Table 3). The remaining five of the 24 patients were from the CAP-O group, where only one (20%) benefited from an intervention from VCE (Table 4, Table 5). Despite
Fig 3. Difference in patients with chronic abdominal pain (CAP). (A) Abnormal findings by referral source. Patients receiving capsule endoscopy for CAP from the community setting had a significantly higher rate of abnormal findings than those referred for capsule studies from a tertiary center. (B) Patients with CAP who also received gastric emptying studies.

doi:10.1371/journal.pone.0126509.g003

Table 2. Sensitivity analysis of patients with chronic abdominal pain and associated symptoms.

| Outcome                        | CAP-A Abnormal VCE | Normal VCE | Total | Obscure Bleeding of GI tract (VCE-GIB-mA) Abnormal VCE | Normal VCE | Total |
|--------------------------------|--------------------|------------|-------|--------------------------------------------------------|------------|-------|
| Spontaneous Resolution         | 3                  | 6          | 9     | 5                                                      | 23         | 27    |
| Intervention Related Resolution| 7                  | 3          | 10    | 24                                                     | 10         | 34    |
| Unresolved                     | 3                  | 21         | 24    | 6                                                      | 18         | 24    |
| Total with Follow Up Data      | 13                 | 30         | 43    | 35                                                     | 51         | 86    |
| No Follow Up Data              | 3                  | 13         | 16    | -                                                      | -          | -     |

Sensitivity analysis of patients with chronic abdominal pain and associated symptoms (CAP-A) and 2:1 age- and gender-matched patients who received a VCE for obscure GI bleeding and had follow up data (VCE-GIB-mA).

doi:10.1371/journal.pone.0126509.t002
the concern for malignancy, none of the 90 patients evaluated for abdominal pain had masses visualized by VCE.

Of the 11 CAP patients who benefitted from an intervention, VCE was abnormal in eight (72.7%). Active Crohn’s disease, enteropathy, and ulcerations were the most common diagnoses following an abnormal study for patients with CAP. Accordingly, the most common interventions based on these diagnoses were implementation of an immune suppression therapy and cessation of NSAID use. Out of the eight cases where VCE led to a change in management, five (62.5%) were due to Crohn’s disease related therapy, in patients with a prior history of the disease.

Table 3. Abnormal findings and outcomes for CAP-A patients.

| Indication         | Sex | Age | Referral Source | Findings                          | Location of Finding | Outcome                |
|--------------------|-----|-----|-----------------|-----------------------------------|---------------------|------------------------|
| CAP + Melena       | F   | 47  | Community       | Abnormal Emptying Diffuse erythema| Stomach & Terminal ileum | CAP Unresolved         |
| CAP + N/V          | F   | 63  | Community       | Patchy erythema                  | Terminal ileum      | CAP unresolved          |
| CAP + Anemia       | F   | 46  | Tertiary Center | Ulceration, narrowing            | Jejunum & Early ileum | CAP Unresolved          |
| CAP + Hematochezia | F   | 25  | Tertiary Center | Ulcerations                      | Jejunum & Ileocecal junction | CAP Unresolved         |
| CAP + Anemia       | F   | 42  | Community       | Gastropathy, ulcers, erosions     | Stomach & Jejunum   | Loss to follow up       |
| CAP + N/V          | F   | 32  | Tertiary Center | Erythema /duodenitis             | Duodenum & Early Jejunum | Loss to follow up       |
| CAP + Diarrhea     | M   | 25  | Community       | Polyps                           | Distal ileum        | Resolved after intervention |
| CAP + PMH of Crohn’s | F | 30  | Community       | Ulceration, nodularity           | Jejunum             | Resolved after intervention |
| CAP + PMH of Crohn’s | F | 52  | Community       | Inflammation, ulceration         | Ileocolonic junction | Resolved after intervention |
| CAP + PMH of Crohn’s | M | 71  | Community       | Erythema and inflammation        | Distal ileum        | Resolved after intervention |
| CAP + PMH Crohn’s + Weight loss | M | 24  | Tertiary Center | Stricture                        | Jejunum             | Resolved after intervention |
| CAP + PMH of Crohn’s | M | 71  | Tertiary Center | Inflamed/ulcerated stricture     | Jejunum & ileum     | Resolved after intervention |
| CAP + Melena       | F   | 32  | Tertiary Center | Ulcers, erythema and inflammation | Stomach & ileum     | Resolved after intervention |
| CAP + N/V          | F   | 44  | Community       | Nodular mucosa and erythema      | Jejunum             | Spontaneous resolution  |
| CAP + N/V          | M   | 26  | Community       | Erythema                         | Terminal ileum      | Spontaneous resolution  |
| CAP + Diarrhea     | M   | 29  | Community       | Erythema/edema                   | Jejunum             | Spontaneous resolution  |

doi:10.1371/journal.pone.0126509.t003

Table 4. Sensitivity Analysis for chronic abdominal pain without associated symptoms.

| Outcome                 | CAP-O | Obscure Bleeding of GI tract (VCE-GIB-mO) |
|-------------------------|-------|------------------------------------------|
|                         | Abnormal VCE | Normal VCE | Total | Abnormal VCE | Normal VCE | Total |
| Spontaneous Resolution  | 1     | 3          | 4     | 3            | 11         | 14    |
| Intervention Related Resolution | 1 | 0        | 1     | 10           | 4          | 14    |
| Unresolved              | 1     | 12         | 13    | 2            | 6          | 8     |
| Total with Follow Up Data | 3    | 15        | 18    | 15           | 21         | 36    |
| No Follow Up            | 3     | 10         | 13    | -            | -          | -     |

Sensitivity analysis of patients with chronic abdominal pain and no other symptoms (CAP-O) and 2:1 age- and gender-matched patients who received a VCE for obscure GI bleeding who had follow up data (VCE-GIB-mO).

doi:10.1371/journal.pone.0126509.t004
In addition, there were three patients who resolved after an intervention, but had a normal VCE. One patient resolved pain with a change in diet. The CAP of the other two resolved with marijuana cessation.

Sensitivity Analysis

In order to determine whether the abnormal finding rate and the frequency of interventions that improved outcomes were similar between patients who received VCE for CAP and patients who received VCE for an indication with strong evidence (i.e., obscure gastrointestinal bleed), we performed a sensitivity analysis. We decided to use VCE-GIB population since the evidence is strong that these patients would benefit from a VCE, and its indication clearly stated in society practice guidelines [10]. In the initial analysis, 86 patients who received VCE for obscure gastrointestinal bleed (VCE-GIB-mA) were randomly selected to correspond with the 43 patients we had follow up data for. These patients were age- and gender-matched to the patients in the CAP-A group (see methods for selection process). In the VCE-GIB-mA group, we found an abnormal finding rate that was not significantly different than that of the CAP-A cohort (40.7% v 27.1%, p = 0.32, respectively, Fig 4A). We repeated this analysis for the CAP-O cohort by age- and gender-matching VCE-GIB-mO patients whom we had follow up data for in a 2:1 ratio (total of 36 patients). Fifteen out of the 36 cases (41.7%) had abnormal findings on VCE study. The CAP-O cohort had significantly lower rate of abnormal findings than the VCE-GIB-mO patients (19.7% v 41.7%, p = 0.009, respectively, Fig 4B).

We found differing results when analyzing patients with successful interventions as a result of VCE. Twenty four out of the 86 cases (27.9%) experienced a change in management or intervention due to the abnormal VCE study. This success rate was significantly different to that of CAP-A patients (11.8% v 27.9%, p = 0.03, respectively, Table 2, Fig 4C). Of the 36 patients in the VCE-GIB-mO cohort, 10 out of the 36 cases (27.8%) experienced a change in management or intervention due to the abnormal VCE study. The successful intervention rate for CAP-O was significantly less than that of the matched VCE-GIB-mO cohort (3.2% v 27.8%, p = 0.008, respectively, Table 4, Fig 4D).

Since many of the CAP-A patients who benefited from an intervention subsequent to VCE were Crohn’s disease patients, we decided to analyze them separately. When the Crohn’s disease patients were removed from analysis, there was still no significant difference between CAP-A and VCE-GIB-mA cohorts in percentage with abnormal findings (23.9% v 40.0%, p = 0.12, respectively, S2 Fig). However, the percentage of CAP-A patients with a successful intervention decreased significantly (4.4% v 23.3%, p = 0.004). Finally, we compared patients with CAP and a history of Crohn’s disease to their matched cohort (VCE-GIB-mCD). In patients with CAP and

Table 5. Abnormal findings and outcomes for chronic abdominal pain (without other symptoms).

| Indication | Sex | Age | Referral Source | Findings | Location of Finding | Outcome |
|------------|-----|-----|-----------------|----------|--------------------|---------|
| CAP Only   | F   | 46  | Community       | Erythematous spots | Jejunum              | CAP unresolved |
| CAP Only   | F   | 52  | Community       | Ulcers    | Distal Jejunum     | Loss to follow up |
| CAP Only   | F   | 56  | Community       | White polypoid protrusion (cyst vs polyp vs lymphoid tissue) | Early Jejunum | Loss to follow up |
| CAP Only   | M   | 36  | Community       | Enteritis  | Early Jejunum      | Loss to follow up |
| CAP Only   | M   | 62  | Community       | Stricture with ulceration | Proximal ileum | Resolved after Intervention |
| CAP Only   | F   | 36  | Community       | Ulcerations, Erythema | Ileum               | Spontaneous resolution |

doi:10.1371/journal.pone.0126509.t005
a history of Crohn’s, there was no significant difference in successful intervention (38.5% v 38.4%, CAP with Crohn’s v VCE-GIB-mCD, p = 1.0, respectively, S3 Fig).

**Discussion**

Patients with CAP, as a whole, were less likely to have abnormal findings on VCE when compared to those who received VCE for other indications. Importantly, no small bowel malignancies were identified in the CAP group. However, when the patients were subdivided into those who had CAP as their sole symptom (CAP-O) versus those who had associated symptoms (CAP-A),
differences started to emerge. VCE rarely was helpful in the CAP-O group. The diagnostic yield of the exam was low. Furthermore, it was unlikely that an abnormal finding would lead to a change in management of the patient’s disease. However, for the CAP-A group, the abnormal finding rate was not different from VCE performed for other indications. Moreover, some patients benefitted since VCE changed management at a rate equivalent compared to VCE performed for gastrointestinal bleeding, which is a well-established indication and part of the AGA and ASGE guidelines for evaluation of obscure gastrointestinal bleeding.

Previous retrospective studies of VCE for CAP have reported various diagnostic rates. However, whether these rates are high or low, or worth the cost of performing the diagnostic test became a matter of debate. Our study contributes to the understanding the role of VCE in CAP in two ways. It first shows whether a higher abnormal finding rate actually translated to clinically meaningful interventions. Furthermore it shows how this rate compares to an age- and gender-matched cohort from the same healthcare system that receives the VCE for a well-established indication. Our study shows that VCE is a useful diagnostic tool that can help provide a possible etiology of CAP in patients with associated symptoms. However, a change in management from information provided by VCE is likely to be limited to those with a history of Crohn’s disease. Despite the presence of strictures and risk of capsule retention, our study did not report any complications from VCE in Crohn’s disease patients. This is consistent with previous literature that shows VCE has significant clinical impact in IBD [11–14].

In our study, VCE yielded abnormal findings in 24.4% of patients with CAP. This rate is consistent with a recent systematic review regarding CAP and VCE, which reported a rate of 20.9% [9]. Out of the 21 studies selected in their review, diagnostic rate varied from 4–44%, with over 12% standard deviation from an unadjusted mean of 22%. The usefulness of VCE in patients with CAP remained dubious due to drastically varying diagnostic yields and other significant limitations in the literature evaluated. This study addresses many of those limitations and also offers a simple classification by which physicians can classify a patient with CAP. It also provides an explanation of why such high variance in diagnostic yield exists since these studies may have had different proportion of CAP-O and CAP-A patients. In CAP-O patients, VCE has a lower diagnostic yield and is much less likely to lead to an intervention that improves outcomes. However, in CAP-A patients, VCE could be a useful tool, providing relevant findings at a rate similar to that of age- and gender-matched patients who were getting VCE for gastrointestinal bleeding. However, even in this group, these findings did not translate to successful interventions. The only group that seemed to benefit were those with Crohn’s disease.

Although the abnormal finding rate and successful intervention percentage may seem low for the VCE-GIB groups, this is likely due to the younger age of patients in this age-matched cohort. Furthermore, as an open referral center, many of the patients with gastrointestinal bleeding had their VCE study performed many months after first referral since patients needed prior insurance approval before the procedure. Previous studies have shown that prolonged time between referral and VCE reduces diagnostic yield in patients with obscure GI bleed [15, 16]. Despite this, we believe our matched gastrointestinal bleeding cohorts remain an adequate measure of comparison on VCE impact, given that our cohorts match the majority of literature regarding outcome and overall clinical impact [16–19].

This study has the largest cohort of patients from the United States, however, other larger studies exist from Asia. The various homogenous populations within Asia have differing prevalence of gastrointestinal disease and infectious disease, which may significantly alter predictors of an abnormal study or the average rate of abnormal findings [20]. Furthermore, significant differences in prevalence and etiology of chronic abdominal pain exist in international states compared to the United States [17, 21]. Despite these differences, the largest study to date (n = 243)
reported a yield (23%) quite similar to our own [22]. This prospective study involved nine tertiary institutions in China. The second largest CAP study, a retrospective study from Korea (n = 110) reported a yield of 17.3% [23]. Neither study reported differences in yield based upon clinical presentation nor whether VCEs changed management or improved outcomes. The larger studies were primarily prospectively evaluating the usefulness of VCE at their respective institutions, all of which were tertiary referral centers. This study is the first to report differences in CAP management and treatment following VCE.

Smaller studies have reported a much greater variation in yield. A multicenter study from Greece (n = 72) reported an overall diagnostic yield of 44.4% for CAP, while another study (n = 64) reported an overall diagnostic yield of 9% [24, 25]. Other studies have reported a diagnostic yield of 36–40% (n = 50) and 6.3% (n = 16) [26, 27]. Hence, our studies abnormal finding yield falls within the large range reported in previous literature. Moreover, it confirms previous reports that the diagnostic yield of VCE improves when CAP is present with other associated symptoms [23–25].

Over the last several years, enterography studies of the small bowel (either by magnetic resonance [MR] or by computed tomography [CT]) have increasingly been used as diagnostic tools in the evaluation of patients with suspected or established IBD [28, 29]. A significant portion of our studies were performed prior to this trend and enterography is a commonly used alternative to VCE in patients with IBD. Whether they are superior to VCE in diagnosis is debatable and outside the scope of this study [30–35]. The role of enterography and any potential advantages in its evaluation of CAP remains unclear. Meanwhile, VCE has proven a cost effective, highly sensitive, and clinically useful modality in the evaluation of IBD that offers a diagnostic advantage in mild to moderate disease activity in Crohn’s disease where mild tissue inflammation and shallow ulcerations may not be obvious on enterography [32, 34, 36–39].

The rate of abnormal findings in this study is similar to that of other diagnostic modalities utilized the evaluation of CAP as well. Colonoscopy and esophagogastroduodenoscopy (EGD) have been reported to have a diagnostic yield of 24%, and 23–46%, respectively. [40, 41]. Unlike EGDS and colonoscopies where ancillary findings such as abnormal imaging and/or serum biomarkers increase the incidence of abnormal findings, yield of VCE in patients with CAP did not improve with the incorporation of these factors. Abnormal blood biomarkers such as ESR, CRP, ANA, and leukocytosis were not associated with study abnormalities in any patient subset. Hence, no serum biomarker measures that were helpful in identifying a subgroup of patients with CAP who may benefit from VCE.

Community physicians were referring the majority of the CAP-O population. Although we presumed that those in our tertiary care center were more likely to have abnormal findings due to the complexity of their cases, community gastroenterologists were more likely to refer patients that would eventually have an abnormal study. Furthermore, the increased use of gastric emptying study in the patients in the CAP-O group further confirms the suspicion that there is an over utility of diagnostic tests in the CAP population.

VCE is a low risk diagnostic modality for visualizing the gastrointestinal tract. As such, it is increasingly being utilized for patients with potential abdominal sources of disease. However, the studies are expensive, require skilled monitoring and review, and present some risk for complications associated with capsule retention. Although none of our patients with CAP had a complication from VCE, capsules can rarely cause an obstruction requiring a surgical intervention. CAP without other associated symptoms appears to be a low yield indication for VCE given a low rate of abnormal findings, and lower likelihood that an abnormal finding could lead to an intervention. In addition, no malignancies of the small bowel were identified. More than a third of patients with CAP had no associated symptoms. This finding is particularly useful for healthcare costs. However, larger, multi-center studies would still be helpful to confidently evaluate whether
VCE is consistently helpful in patients with associated symptoms or a worrisome past medical history.

While this large retrospective study sampled a diverse population of patients from throughout San Diego and Imperial Counties, it has its limitations. We did not have follow up data for all of the patients who received VCE for CAP. Furthermore, chart review was limited by the completeness of physician documentation and did not allow for interpretation of intent, unless explicitly stated. Hence, we were unable to quantify any benefit of a negative study. Also, we acknowledge the inability of this study to comment on the characteristics of patients who had abdominal pain, but were not referred to the UCSD endoscopy center for a VCE. Such patients may represent those with less impressive clinical presentations or lower pre-test probability of disease.

In conclusion, the role of VCE in CAP remains complex. Although VCE is unlikely to be beneficial in patients without associated symptoms, it may play a role in those who have associated symptoms. In this study, VCE not only had an abnormal finding rate that was comparable to upper and lower endoscopy, but also was likely to lead to altered disease management. More prospective studies are needed to better understand the role that VCE can play in CAP.

Supporting Information

S1 Fig. Flowchart with classification of patients from the control group; patients evaluated for reasons other than chronic abdominal pain group. Sub-group classification based on similar presentation. Abbreviations: GI (gastrointestinal), HHT (heredity hemorrhagic telangiectasia), Hx (history), FAP (familial adenomatous polyposis).

S2 Fig. Abnormal Finding Rate if Crohn’s Disease Patients Are Excluded. Abnormal finding rate comparison between patients with CAP-A, but excluding patients with a history of Crohn’s disease (CAP-A without CD) and their 2:1 age- and gender-matched comparison group as previously described, without Crohn’s disease matched patients (VCE-GIB-mA-CD).

S3 Fig. Inclusion and Exclusion of Crohn’s Disease Patients on Treatment Outcomes. CAP with CD, CAP-A without CD, and their 2:1 age, gender matched control group (VCE-GIB-mCD, VCE-GIB-mA-CD, respectively). ** represents a p-value of < 0.01.

S1 Table. Comparison of Inflammatory markers (not available for many patients) versus abnormal findings. Abbreviations: ESR: Erythrocyte sedimentation rate. CRP: C Reactive Protein.

S2 Table. Abnormal findings and outcomes for patients with chronic abdominal pain (CAP) and Crohn’s disease.

S3 Table. Sensitivity analysis of patients with Crohn’s Disease (CD) and chronic abdominal pain (CA) compared to both patients with CAP and associated symptoms without Crohn’s Disease (CAP-A without CD), the 2:1 gender and aged matched patients for CAP-A without CD in patients who received a VCE for gastrointestinal bleeding and had follow up data (VCE-GIB-mA-CD).
Acknowledgments

The authors would like to thank Divya Chandrasekar who helped form an earlier database of video capsule endoscopies which was used to initiate this study. We would also like to thank Dr. Thomas J. Savides and Dr. Samuel Ho for critical review of earlier drafts and Dr. William Sandborn for critical discussion.

Author Contributions

Conceived and designed the experiments: DK AZ. Performed the experiments: JE KK AZ. Analyzed the data: JE AZ. Wrote the paper: JE AZ.

References

1. Quigley EM, Locke GR, Mueller-Lissner S, Paulo LG, Tytgat GN, Heilrich I, et al. Prevalence and management of abdominal cramping and pain: a multinational survey. Alimentary pharmacology & therapeutics. 2006; 24(2):411–9. doi:10.1111/j.1365-2036.2006.02989.x PMID: 16842469.

2. Sandler RS, Stewart WF, Liberman JN, Ricci JA, Zorich NL. Abdominal pain, bloating, and diarrhea in the United States: prevalence and impact. Digestive diseases and sciences. 2000; 45(6):1166–71. PMID: 10877233.

3. Townsend CO, Sletten CD, Bruce BK, Rome JD, Luedtke CA, Hodgson JE. Physical and emotional functioning of adult patients with chronic abdominal pain: comparison with patients with chronic back pain. The journal of pain: official journal of the American Pain Society. 2005; 6(2):75–83. doi:10.1016/j.jpain.2004.10.009 PMID: 15694873.

4. Hulisz D. The burden of illness of irritable bowel syndrome: current challenges and hope for the future. Journal of managed care pharmacy: JMCP. 2004; 10(4):299–309. PMID: 15298528.

5. Middleton KR, Hing E. National Hospital Ambulatory Medical Care Survey: 2004 outpatient department summary. Advance data. 2006;(373): 1–27. PMID: 16841784.

6. Myer PA, Mannalithara A, Singh G, Singh G, Pasricha PJ, Ladabaum U. Clinical and economic burden of emergency department visits due to gastrointestinal diseases in the United States. Am J Gastroenterol. 2013; 108(9):1496–507. doi: 10.1038/ajg.2013.199 PMID: 23857475.

7. Makins R, Blanshard C. Guidelines for capsule endoscopy: diagnoses will be missed. Alimentary pharmacology & therapeutics. 2006; 24(2):293–7. doi: 10.1111/j.1365-2036.2006.02991.x PMID: 16638706.

8. Qvigstad G, Hatlen-Rebhan P, Brenna E, Waldum HL. Capsule endoscopy in clinical routine in patients with suspected disease of the small intestine: a 2-year prospective study. Scandinavian journal of gastroenterology. 2006; 41(5):614–8. doi: 10.1080/0036552050035159 PMID: 16638706.

9. Xue M, Chen X, Shi L, Si J, Wang L, Chen S. Small-bowel capsule endoscopy in patients with unexplained chronic abdominal pain: a systematic review. Gastrointestinal endoscopy. 2015; 81(1):186–93. doi: 10.1016/j.gie.2014.04.062 PMID: 25012561.

10. Raju GS, Gerson L, Das A, Lewis B. American Gastroenterological A. American Gastroenterological Association (AGA) Institute technical review on obscure gastrointestinal bleeding. Gastroenterology. 2007; 133(5):1697–717. doi: 10.1053/j.gastro.2007.06.007 PMID: 17953812.

11. Cheifetz AS, Lewis BS. Capsule endoscopy retention: is it a complication? J Clin Gastroenterol. 2006; 40(8):688–91. Epub 2006/08/31. PMID: 16940879.

12. Kopylov U, Seidman EG. Role of capsule endoscopy in inflammatory bowel disease. World J Gastroenterol. 2014; 20(5):1155–64. Epub 2014/02/28. doi: 10.3748/wjg.v20.i5.1155 PMID: 24574792; PubMed Central PMCID: PMC3921500.

13. Yamagami H, Watanabe K, Kamata N, Sogawa M, Arakawa T. Small bowel capsule endoscopy in inflammatory bowel disease. Clin Endosc. 2013; 46(4):321–6. Epub 2013/08/22. doi: 10.5946/ce.2013.46.4.321 PMID: 23964326; PubMed Central PMCID: PMC3746136.

14. Long MD, Barnes E, Isaacs K, Morgan D, Herfarth HH. Impact of capsule endoscopy on management of inflammatory bowel disease: a single tertiary care center experience. Inflamm Bowel Dis. 2011; 17(9):1855–62. Epub 2011/08/11. doi: 10.1002/ibd.21571 PMID: 21830264; PubMed Central PMCID: PMC3116881.

15. Esaki M, Matsumoto T, Yada S, Yanaru-Fujisawa R, Kudo T, Yanai S, et al. Factors associated with the clinical impact of capsule endoscopy in patients with overt obscure gastrointestinal bleeding. Dig Dis Sci. 2010; 55(8):2294–301. Epub 2009/12/04. doi: 10.1007/s10620-009-1036-5 PMID: 19957038.
16. Santhakumar C, Liu K. Evaluation and outcomes of patients with obscure gastrointestinal bleeding. World J Gastrointest Pathophysiol. 2014; 5(4):479–86. doi: 10.4291/wjgp.v5.i4.479 PMID: 25400992; PubMed Central PMCID: PMC4231513.

17. Heo HM, Park CH, Lim JS, Lee JH, Kim BK, Cheon JH, et al. The role of capsule endoscopy after negative CT enterography in patients with obscure gastrointestinal bleeding. Eur Radiol. 2012; 22(6):1159–66. Epub 2012/01/25. doi: 10.1007/s00330-011-2374-1 PMID: 22270143.

18. Lai LH, Wong GL, Chow DK, Lau JY, Sung JJ, Leung WK. Long-term follow-up of patients with obscure gastrointestinal bleeding after negative capsule endoscopy. Am J Gastroenterol. 2006; 101(6):1224–8. Epub 2006/06/15. doi: 10.1111/j.1572-0241.2006.00565.x PMID: 16771942.

19. Rondonotti E, Soncini M, Girelli C, Ballardi G, Bianchi G, Brunati S, et al. Small bowel capsule endoscopy in clinical practice: a multicenter 7-year survey. Eur J Gastroenterol Hepatol. 2010; 22(11):1380–6. Epub 2010/02/23. doi: 10.1097/MEG.0b013e3283352ced PMID: 20173646.

20. Ng SC. Inflammatory Bowel Disease in Asia. Gastroenterol Hepatol (N Y). 2013; 9(1):28–30. PMID: 24707231; PubMed Central PMCID: PMC3975975.

21. Katsinelos P, Tziomalos K, Fasoulas K, Paroutoglou G, Koutokosios A, Mimidis K, et al. Can capsule endoscopy be used as a diagnostic tool in the evaluation of nonbleeding indications in daily clinical practice? A prospective study. Med Princ Pract. 2011; 20(4):362–7. doi: 10.1159/000324548 PMID: 21576998.

22. Yang L, Chen Y, Zhang B, Chen C, Yue M, Du J, et al. Increased diagnostic yield of capsule endoscopy in patients with chronic abdominal pain. PLoS one. 2014; 9(1):e87396. doi: 10.1371/journal.pone.0087396 PMID: 24489097; PubMed Central PMCID: PMC3909121.

23. Shim KN, Kim YS, Kim KJ, Kim YH, Kim TI, Do JH, et al. Abdominal pain accompanying weight loss may increase the diagnostic yield of capsule endoscopy: a Korean multicenter study. Scandinavian journal of gastroenterology. 2006; 41(8):983–8. doi: 10.1080/00365520600548974 PMID: 16803698.

24. Fry LC, Carey EJ, Shiff AD, Heigh RI, Sharma VK, Post JK, et al. The yield of capsule endoscopy in patients with abdominal pain or diarrhea. Endoscopy. 2006; 38(5):498–502. doi: 10.1055/s-2006-925340 PMID: 16767586.

25. Katsinelos P, Fasoulas K, Beltsis A, Chatzimavroudis G, Paroutoglou G, Maris T, et al. Diagnostic yield and clinical impact of wireless capsule endoscopy in patients with chronic abdominal pain with or without diarrhea: a Greek multicenter study. European journal of internal medicine. 2011; 22(5):e63–6. doi: 10.1016/j.ejim.2011.06.012 PMID: 21925046.

26. May A, Manner H, Schneider M, Ipsen A, Ell C. Prospective multicenter trial of capsule endoscopy in patients with chronic abdominal pain, diarrhea and other signs and symptoms (CEDAP-Plus Study). Endoscopy. 2007; 39(7):606–12. doi: 10.1055/s-2007-966640 PMID: 17611915.

27. Spada C, Pirozzi GA, Riccioni ME, Iacopini F, Marchese M, Costamagna G. Capsule endoscopy in patients with chronic abdominal pain. Digestive and liver disease: official journal of the Italian Society of Gastroenterology and the Italian Association for the Study of the Liver. 2006; 38(9):696–9. doi: 10.1016/j.dld.2006.05.011 PMID: 16920049.

28. Santillan CS. MR imaging techniques of the bowel. Magnetic resonance imaging clinics of North America. 2014; 22(1):1–11. doi: 10.1016/j.mrc.2013.07.004 PMID: 24238128.

29. Zamboni GA, Rapotopoulos V. CT enterography. Gastrointestinal endoscopy clinics of North America. 2010; 20(2):347–66. doi: 10.1016/j.geic.2010.02.017 PMID: 20451821.

30. Solem CA, Loftus EV Jr., Fletcher JG, Baron TH, Gostout CJ, Petersen BT, et al. Small-bowel imaging in Crohn's disease: a prospective, blinded, 4-way comparison trial. Gastrointestinal endoscopy. 2008; 68(2):255–66. doi: 10.1016/j.gie.2008.02.017 PMID: 18513722.

31. Jensen MD, Nathan T, Rafaeislen SR, Kjeldsen J. Diagnostic accuracy of capsule endoscopy for small bowel Crohn's disease is superior to that of MR enterography or CT enterography. Clinical gastroenterology and hepatology: the official clinical practice journal of the American Gastroenterological Association. 2011; 9(2):124–9. doi: 10.1016/j.cgh.2010.10.019 PMID: 21056699.

32. Hara AK, Leighton JA, Heigh RI, Sharma VK, Silva AC, De Petris G, et al. Crohn disease of the small bowel: preliminary comparison among CT enterography, capsule endoscopy, small-bowel follow-through, and ileoscopy. Radiology. 2006; 238(1):128–34. doi: 10.1148/radiol.2381050296 PMID: 16373764.

33. Triester SL, Leighton JA, Leontiadis GI, Gurudu SR, Fleischer DE, Hara AK, et al. A meta-analysis of the yield of capsule endoscopy compared to other diagnostic modalities in patients with non-stricturing small bowel Crohn's disease. Am J Gastroenterol. 2006; 101(5):954–64. Epub 2006/05/16. doi: 10.1111/j.1572-0241.2006.00506.x PMID: 16636781.

34. Dionisio PM, Gurudu SR, Leighton JA, Leontiadis GI, Fleischer DE, Hara AK, et al. Capsule endoscopy has a significantly higher diagnostic yield in patients with suspected and established small-bowel Crohn's disease: a meta-analysis. Am J Gastroenterol. 2010; 105(6):1240–8; quiz 9. Epub 2009/12/24. doi: 10.1038/ajg.2009.713 PMID: 20029412.
35. Leighton JA, Triester SL, Sharma VK. Capsule endoscopy: a meta-analysis for use with obscure gastrointestinal bleeding and Crohn's disease. Gastrointest Endosc Clin N Am. 2006; 16(2):229–50. Epub 2006/04/29. doi: 10.1016/j.giec.2006.03.004 PMID: 16644453.

36. Goldfarb NI, Pizzi LT, Fuhr JP Jr., Salvador C, Sikirica V, Kornbluth A, et al. Diagnosing Crohn's disease: an economic analysis comparing wireless capsule endoscopy with traditional diagnostic procedures. Dis Manag. 2004; 7(4):292–304. Epub 2005/01/27. doi: 10.1089/dis.2004.7.292 PMID: 15671786.

37. Hamilton MJ. The Valuable Role of Endoscopy in Inflammatory Bowel Disease. Diagnostic and Therapeutic Endoscopy. 2012; 2012. doi: 10.1155/2012/467979.

38. Jewell DM, Neil, Steinhart, Hillary. Challenges in Inflammatory Bowel Disease. 2nd ed: Wiley-Blackwell; 2006. 4 p.

39. Swain P. Wireless capsule endoscopy and Crohn's disease. Gut. 542005, p. 323–6.

40. Morini S, Hassan C, Meucci G, Toldi A, Zullo A, Minoli G. Diagnostic yield of open access colonoscopy according to appropriateness. Gastrointestinal Endoscopy. 2001; 54(2):175–9. doi: 10.1067/mge.2001.116565 PMID: 11474386.

41. Burri E, Manz M, Schroeder P, Froehlich F, Rossi L, Beglinger C, et al. Diagnostic yield of endoscopy in patients with abdominal complaints: incremental value of faecal calprotectin on guidelines of appropriateness. BMC Gastroenterology. 2014; 14(1):57. info: PMID: 24679065.