

Implementing Population Medicine in a Pain Management Practice

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ABSTRACT

Purpose: To document and improve the quality of our chronic pain management using population management methods.

Methods: An analytic registry was developed, and all new patients were enrolled for 12 months. Patient demographics, standardized pain and function measures, and treatments were recorded. Usual care was provided. The registry was used to organize care and analyze management and outcomes.

Results: Of 454 total patients, only 154 (34%) completed a 6-month cycle of care. High no-show rates were documented for follow-up appointments for several reasons. The majority of 6-month completers showed improved pain levels.

Discussion: This quality improvement project identified assessment and care gaps and led to improvements. An ongoing need to improve measures of pain and function was documented.

INTRODUCTION

Managing chronic pain and its underlying causes presents a continuing challenge to health systems, clinicians, patients, and health planners in the United States.¹ Diagnoses and treatments for similar patients vary greatly by individual physician and across specialties. Opioid dependency is but one highly visible problem associated with current approaches to pain management. In fact,

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the fragmented and variable care processes, high costs, and suboptimal outcomes of chronic pain management differ little from those of most chronic diseases.²

Population medicine utilizes a disease registry to identify all patients within a population and to then guide care teams to provide and document necessary care on time. This approach changes the focus of care from one patient at a time to the population as a whole, with individual patient's care provided within this broader context. It allows practices to identify and close care gaps that cannot otherwise be appreciated or addressed. In general, population medicine is proving more effective than traditional approaches for managing chronic diseases.³

We have implemented population medicine methods within our pain management practice in an effort to improve our care and patient outcomes. This report describes our quality improvement project and initial results.

METHODS

Participants

An interventional pain physician (DAB) and a nurse practitioner (BK) conducted this project in a community-based pain management practice at 3 clinic sites in south central Wisconsin. A physician consultant (JTH) provided quality improvement and population medicine coaching. Our processes were developed to support best clinical practices and high clinical utility, and usual treatments were provided.

Procedure

We began by defining a set of standard disease and treatment data that we intended to collect routinely at baseline and each follow-up patient assessment. A data collection sheet was developed to capture this information (Figure 1). Treatments reported were those provided since the last assessment. The Patient Pain and Provider Global Scores (PGS) were reported on 0-10 segmented

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Figure 1. Data Collection Sheet

| Date | Name | Date of Birth | Registry Number |
|---------------------------|---------------------------------|----------------------|------------------------------|
| | | | |
| Parameters | | Treatments | |
| | <i>Patient derived</i> | | <i>Medications</i> |
| VAS- current rating | | Opioids | |
| VAS- maximum rating | | Neuroleptics | |
| Oswestry score | | Non steroidal | |
| | <i>Provider derived</i> | Local anesthetics | |
| Opioid risk score | | Antidepressants | |
| MEDD | | Mood stabilizers | |
| Practitioner Global Score | | | <i>Interventional</i> |
| | <i>Cycle of Care - 6 months</i> | Diagnostic | |
| Initial visit | | Therapeutic | |
| One | | Neuromodulation | |
| Two | | Radiofrequency | |
| Three | | | <i>Psychology Evaluation</i> |
| | | Cognitive therapy | |
| | | Group therapy | |
| | | Individual therapy | |
| | | | <i>Therapy</i> |
| | | Physical therapy | |
| | | Occupational therapy | |

Standardized pain, function, and global measures and current treatments are collected at each patient assessment visit, generally at baseline, 6, 12, and 24 weeks.

Abbreviations: VAS, visual analogue scales; MEDD, Morphine Equivalent Daily Dose.

visual analogue scales (VAS) with 0 to .99 = Controlled, 1 to 3.9 = Low, 4 to 6.9 = Moderate, and 7 to 10 = High levels of pain and disability.⁴ Opioid use was calculated as morphine equivalent daily dose (MEDD) from a standard conversion table, and the Oswestry Disability Index was calculated from a patient-generated questionnaire and segmented into low, moderate, and high levels.^{5,6} The PGS was used to capture the clinician's overall impression based on patient history, other objective patient-derived measures, and examination findings. A PGS has been used widely in clinical trials and for documenting the activity of other chronic diseases⁷ but, to our knowledge, not to capture the provider's overall impression in pain management.

We then determined the intervals at which we wished to follow patients after baseline evaluation and initiation of medical or procedural treatments, recognizing that these would vary in some cases. We assumed that our cycle of care was generally 6 months in duration and that assessments would be performed at 6 weeks, 12 weeks, and 24 weeks.

We next developed a disease population registry in an Excel database, backed up and protected on our practice's HIPAA-

compliant information technology platform. We enrolled each new patient with their identifying information, referral source, date of consultation, and International Classification of Diseases, Ninth Revision (ICD-9) diagnoses. Each patient's clinical data were entered at baseline and at each follow-up assessment. Our analytic registry format is shown in Figure 2.

A nurse practitioner (BK) managed the registry. Data from collection sheets were entered real-time or batched and then entered separately as time permitted. The registry spreadsheet was sorted regularly by encounter dates to identify overdue patients and to analyze other data as indicated in Results. Encounters for each patient were also documented in our electronic medical record (EMR), including scanned data collection sheets.

We did not obtain Institutional Review Board oversight since quality improvement projects are generally exempt, as their purpose is to improve care delivery processes and not to perform research or deviate from usual treatments.^{8,9}

RESULTS

The study population included all new patients seen for initial evaluation during a 12-month interval beginning on January 2, 2014 (N = 520). These patients were referred either by community primary (84%) and specialty (15.5%) physicians or were self-referred (0.5%). Patients were 54% male and 46% female and ranged from 20 to 89 years (median = 57). Sixty-six patients with more than 1 pain problem at enrollment were excluded from this analysis, leaving a total of 454. Initial ICD-9 codes included spine disorders (68%), other musculoskeletal conditions (18%), neuropathies (10%), and a variety of other diagnoses (3.5%).

Only 154 (34%) of these 454 enrollees had both baseline and 6-month assessments. An additional 146 (32%) did not keep their first scheduled follow-up appointment with major reasons being lack of insurance coverage and patient decisions to decline recommended care. The remaining 154 (34%) completed specialty pain management in less than 6 months. They were returned to their referring physician for medication management, referred to other specialists, continued in rehabilitation, and/or had resolved their pain problem. Many of this latter cohort did not have a discharge visit and assessment.

A variety of interventional procedures were performed on 170

In conclusion, this population medicine project has improved our interventional pain management practice. We hope our experiences will encourage others to adopt population medicine approaches and standardized measures of patients' status and outcomes, not only in pain management practices, but also within other specialties and broader health systems. Documenting and improving care and outcomes for chronic disease populations are critical to increasing the value of care and overcoming barriers to payment for effective services.

Funding/Support: None declared.

Financial Disclosures: None declared.

Planners/Reviewers: The planner and reviewers for this journal CME activity have no financial relationship to disclose.

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