Dear Mr. Spencer

The Division of Health Improvement Quality Management Bureau has completed a focused survey of the service identified above. The purpose of the survey was to determine compliance with federal and state standards; to assure the health, safety, and welfare of individuals receiving services through the Developmental Disabilities Waiver; and to identify opportunities for improvement. The specific focus of the survey was to determine compliance regarding Medical Oversight and Direct Service Personnel training.

Plan of Correction:
The attached Report of Findings identifies deficiencies found during your agency’s survey. You are required to complete and implement a Plan of Correction (POC). Please submit your agency’s Plan of Correction (POC) in the space on the two right columns of the Report of Findings. See attachment A for additional guidance in completing the POC. The response is due to the parties below within 10 working days of the receipt of this letter:

1. Quality Management Bureau, Attention: Plan of Correction Coordinator
   5301 Central Ave. NE Suite 400 Albuquerque, NM 87108

2. Developmental Disabilities Supports Division Regional Office for region of service surveyed.

Upon notification from QMB that your Plan of Correction has been approved, you must implement all remedies and corrective actions within 45 working days. If your plan of correction is denied, you must resubmit a revised plan ASAP for approval. All remedies must still be completed within 45 working days of the original submission.

“Assuring safety and quality of care in New Mexico’s health facilities and community-based programs.”

David Rodriguez, Division Director • Division of Health Improvement
Division of Health Improvement • Quality Management Bureau • 5301 Central Ave NE • Suite 400 • Albuquerque, New Mexico 87108
(505) 222-8623 • FAX: (505) 841-5815


Report #: Q10.01.D1808.SE.001.FCD.01
Failure to submit, complete or implement your POC within the required time frames will result in the imposition of a $200 per day Civil Monetary Penalty until it is received, completed and/or implemented.

**Request for Informal Reconsideration of Findings (IRF):**
If you disagree with a determination of noncompliance (finding) you have 10 working days upon receipt of this notice to request an IRF. Submit your request for an IRF in writing to:

QMB Deputy Bureau Chief  
5301 Central Ave NE Suite #400  
Albuquerque, NM 87108  
Attention: IRF request

A request for an IRF will not delay the implementation of your Plan of Correction which must be completed within 45 working days. Providers may not appeal the nature or interpretation of the standard or regulation, the team composition, sampling methodology or the Scope and Severity of the finding.

If the IRF approves the change or removal of a finding, you will be advised of any changes.

This IRF process is separate and apart from the Informal Dispute Resolution (IDR) and Fair Hearing Process for Sanctions from DOH.

Please call the Team Leader at 505-690-4693, if you have questions about the survey or the report. Thank you for your cooperation and for the work you perform.

Sincerely,

**Deb Russell, BS**

Deb Russell  
Team Lead/Healthcare Surveyor  
Division of Health Improvement  
Quality Management Bureau
Survey Process Employed:

Entrance Conference Date: August 24, 2009

Present:

**ENMRSH, Inc.**
Damian Houfek, Vice President

**DOH/DHI/QMB**
Deb Russell, BS, Team Lead/Healthcare Surveyor
Cynthia Nielsen, RN, MSN, Healthcare Surveyor
Florie Alire, RN, Healthcare Surveyor

Exit Conference Date: August 26, 2009

Present:

**ENMRSH, Inc.**
Robert Spencer, President/CEO
Damian Houfek, Vice President
Kathy Lynch, RN

**DOH/DHI/QMB**
Deb Russell, BS, Team Lead/Healthcare Surveyor
Cynthia Nielsen, RN, MSN, Healthcare Surveyor
Florie Alire, RN, Healthcare Surveyor

**DDSD - Southeast Regional Office**
Jon Hellebust, Regional Program Manager
Sandra Renteria, RN

Administrative Locations Visited Number: 1

Total Sample Size Number: 9

- 3 - Jackson Class Members
- 6 - Non-Jackson Class Members
- 6 - Supported Living
- 2 - Independent Living
- 6 - Adult Habilitation
- 2 - Community Access
- 3 - Supported Employment
- 1 - No Longer Receiving Services

Records Reviewed (Persons Served) Number: 9

Administrative Files Reviewed

- Medical Records
- Incident Management Records
- Personnel Files
- Training Records
- Agency Policy and Procedure
- Nursing personnel files

CC: Distribution List: DOH - Division of Health Improvement
DOH - Developmental Disabilities Supports Division
DOH - Office of Internal Audit
HSD - Medical Assistance Division
Provider Instructions for Completing the QMB Plan of Correction (POC) Process

- After a QMB Quality Review, your Survey Report will be sent to you via certified mail. You may request that it also be sent to you electronically by calling George Perrault, Plan of Correction Coordinator at 505-222-8624.
- Within 10 business days of the date you received your survey report, you must develop and send your Plan of Correction response to the QMB office. (Providers who do not pick up their mail will be referred to the Internal Review Committee [IRC]).
- For each Deficiency in your Survey Report, include specific information about HOW you will correct each Deficiency, WHO will fix each Deficiency (“Responsible Party”), and by WHEN (“Date Due”).
- Your POC must not only address HOW, WHO and WHEN each Deficiency will be corrected, but must also address overall systemic issues to prevent the Deficiency from reoccurring, i.e., Quality Assurance (QA). Your description of your QA must include specifics about your self-auditing processes, such as HOW OFTEN you will self-audit, WHO will do it, and WHAT FORMS will be used.
- Corrective actions should be incorporated into your agency’s Quality Assurance/Quality Improvement policies and procedures.
- You may send your POC response electronically to George.Perrault@state.nm.us, by fax (505-841-5815), or by postal mail.
- Do not send supporting documentation to QMB until after your POC has been approved by QMB.
- QMB will notify you if your POC has been “Approved” or “Denied”.
- Whether your POC is “Approved” or “Denied”, you have a maximum of 45 business days to correct all survey Deficiencies from the date of receipt of your Survey Report. If your POC is “Denied” it must be revised and resubmitted ASAP, as the 45 working day limit is in effect. Providers whose revised POC is denied will be referred to the IRC.
- The POC must be completed on the official QMB Survey Report and Plan of Correction Form, unless approved in advance by the POC Coordinator.
- The following Deficiencies must be corrected within the deadlines below (after receipt of your Survey Report):
  - CCHS and EAR: 10 working days
  - Medication errors: 10 working days
  - IMS system/training: 20 working days
  - ISP related documentation: 30 working days
  - DDSD Training: 45 working days
- If you have questions about the POC process, call the QMB POC Coordinator, George Perrault at 505-222-8624 for assistance.
- For Technical Assistance (TA) in developing or implementing your POC, contact your local DDSD Regional Office.
- Once your POC has been approved by QMB, the POC may not be altered or the dates changed.
- Requests for an extension or modification of your POC (post approval) must be made in writing and submitted to the POC Coordinator at QMB, and are approved on a case-by-case basis.
- When submitting supporting documentation, organize your documents by Tag #s, and annotate or label each document using Individual #s.
- Do not submit original documents, hard copies or scanned and electronically submitted copies are fine. Originals must be maintained in the agency/client file(s) as per DDSD Standards.
- Failure to submit, complete or implement your POC within the required timeframes will result in a referral to the IRC and the possible imposition of a $200 per day Civil Monetary Penalty until it is received, completed and/or implemented.
QMB Scope and Severity Matrix of survey results

Each deficiency in your Report of Findings is scored on a Scope and Severity Scale. The culmination of each deficiency’s Scope and Severity is used to determine degree of compliance to standards and regulations and level of QMB Certification.

<table>
<thead>
<tr>
<th>SCOPE</th>
<th>Isolated 01% - 15%</th>
<th>Pattern 16% - 79%</th>
<th>Widespread 80% - 100%</th>
</tr>
</thead>
<tbody>
<tr>
<td>SEVERITY</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>High Impact</td>
<td>Immediate Jeopardy to individual health and or safety</td>
<td>J.</td>
<td>K.</td>
</tr>
<tr>
<td>Medium Impact</td>
<td>No Actual Harm Potential for more than minimal harm</td>
<td>D.</td>
<td>E.</td>
</tr>
<tr>
<td>Low Impact</td>
<td>No Actual Harm Minimal potential for harm.</td>
<td>A.</td>
<td>B.</td>
</tr>
<tr>
<td></td>
<td>Low Impact Severity: (Blue)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Medium Impact Severity: (Tan)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Medium level findings have a potential for harm to an individual. Providers that have no findings above a “F” level and/or no more than two F level findings and no F level Conditions of Participation may receive a “Merit” Certification approval rating from QMB.

High Impact Severity: (Green or Yellow)
High level findings are when harm to an individual has occurred. Providers that have no findings above “I” level may only receive a “Standard” Approval rating from QMB and will be referred to the IRC.

High Impact Severity: (Yellow)
“J, K, and L” Level findings:
This is a finding of Immediate Jeopardy. If a provider is found to have “I” level findings or higher, with an outcome of Immediate Jeopardy, including repeat findings or Conditions of Participation they will be referred to the Internal Review Committee.
Guidelines for the Provider
Informal Reconsideration of Finding (IRF) Process

Introduction:
Throughout the process, surveyors are openly communicating with providers. Open communication means that surveyors have clarified issues and/or requested missing information before completing the review. Regardless, there may still be instances where the provider disagrees with a specific finding.

To informally dispute a finding the provider must request in writing an Informal Reconsideration of the Finding (IRF) to the QMB Deputy Bureau Chief within 10 working days of receipt of the final report.

The written request for an IRF must be completed on the QMB Request for Informal Reconsideration of Finding Form (available on the QMB website: http://dhi.health.state.nm.us/qmb) and must specify in detail the request for reconsideration and why the finding is inaccurate. The IRF request must include all supporting documentation or evidence that was not previously reviewed during the survey process.

The following limitations apply to the IRF process:

- The request for an IRF and all supporting evidence must be received in 10 days.
- Findings based on evidence requested during the survey and not provided may not be subject to reconsideration.
- The supporting documentation must be new evidence not previously reviewed by the survey team.
- Providers must continue to complete their plan of correction during the IRF process.
- Providers may not request an IRF to challenge the Scope and Severity of a finding.
- Providers may not request an IRF to challenge the sampling methodology.
- Providers may not request an IRF based on disagreement with the nature of the standard or regulation.
- Providers may not request an IRF to challenge the team composition.
- Providers may not request an IRF to challenge the QMB Quality Approval Rating and the length of their DDSD provider contract.

A Provider forfeits the right to an IRF if the request is not made within 10 working days of receiving the report and does not include all supporting documentation or evidence to show compliance with the standards and regulations.

QMB has 30 working days to complete the review and notify the provider of the decision. The request will be reviewed by the IRF committee. The Provider will be notified in writing of the ruling, no face to face meeting will be conducted.

When a Provider requests that a finding be reconsidered, it does not stop or delay the Plan of Correction process. Providers must continue to complete the Plan of Correction, including the finding in dispute regardless of the IRF status. If a finding is successfully reconsidered, it will be noted and will be removed or modified from the report. It should be noted that in some cases a Plan of Correction may be completed prior to the IRF process being completed. The provider will be notified in writing on the decisions of the IRF committee.

Administrative Review Process:
If a Provider desires to challenge the decision of the IRF committee they may request an Administrative Review by the DHI and DDSD Director. The Request must be made in writing to the QMB Bureau Chief and received within 5 days of notification from the IRF decision.

Regarding IRC Sanctions:
The Informal Reconsideration of the Finding process is a separate process specific to QMB Survey Findings and should not be confused with any process associated with IRC Sanctions.

If a Provider desires to Dispute or Appeal an IRC Sanction that is a separate and different process. Providers may choose the Informal Dispute Resolution Process or the Formal Medicaid Fair Hearing Process to dispute or appeal IRC sanctions, please refer to the DOH Sanction policy and section 39 of the provider contract agreement.
**Agency:** ENMRSH, Inc. - Southeast Region  
**Program:** Developmental Disabilities Waiver  
**Service:** Community Living (Supported Living & Independent Living) & Community Inclusion (Adult Habilitation, Community Access & Supported Employment)  
**Monitoring Type:** Focused Survey  
**Date of Survey:** August 24 – 26, 2009

<table>
<thead>
<tr>
<th>Statute</th>
<th>Deficiency</th>
<th>Agency Plan of Correction and Responsible Party</th>
<th>Date Due</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tag # 1A09 Medication Delivery (MAR) - Routine Medication</td>
<td>Scope and Severity Rating: E</td>
<td>Based on record review, 5 of 9 individuals had Medication Administration Records, which contained missing medications entries and/or other errors. Medication Administration Records (MAR) were reviewed for the months of May, June &amp; July 2009. Individual #1 June 2009 As indicated by the Medication Administration Records the individual is to take Naprosyn 500mg (2 times daily). According to the Physician’s Orders written May 19, 2009, Naprosyn 500mg is to be taken 2 times daily for 2 weeks, then PRN. Medication Administration Record &amp; Physician’s Orders do not match. July 2009 As indicated by the Medication Administration Records the individual is to take Naprosyn 500mg (2 times daily). According to the Physician’s Orders written May 19, 2009, Naprosyn 500mg is to be taken 2 times daily for 2 weeks, then PRN. Medication Administration Record &amp; Physician’s Orders do not match.</td>
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</table>
and generic name of the medication, diagnosis for which the medication is prescribed;
(b) Prescribed dosage, frequency and method/route of administration, times and dates of administration;
(c) Initials of the individual administering or assisting with the medication;
(d) Explanation of any medication irregularity;
(e) Documentation of any allergic reaction or adverse medication effect; and
(f) For PRN medication, an explanation for the use of the PRN medication shall include observable signs/symptoms or circumstances in which the medication is to be used, and documentation of effectiveness of PRN medication administered.

3) The Provider Agency shall also maintain a signature page that designates the full name that corresponds to each initial used to document administered or assisted delivery of each dose;

4) MARs are not required for individuals participating in Independent Living who self-administer their own medications;

5) Information from the prescribing pharmacy regarding medications shall be kept in the home and community inclusion service locations and shall include the expected desired outcomes of administering the medication, signs and symptoms of adverse events and interactions with other medications;

NMAC 16.19.11.8 MINIMUM STANDARDS:
A. MINIMUM STANDARDS FOR THE DISTRIBUTION, STORAGE, HANDLING AND RECORD KEEPING OF DRUGS:

(d) The facility shall have a Medication Administration Record (MAR) documenting medication administered to residents, including over-the-counter medications. This documentation shall include:

- Records the individual is to take Eskalith – Lithium 300mg (2 times daily). According to the Physician’s Orders 5/7/2009, Eskalith – Lithium 300mg is to be taken 1 time daily Medication Administration Record & Physician’s Orders do not match.

July 2009
As indicated by the Medication Administration Records the individual is to take Eskalith – Lithium 300mg (2 times daily). According to the Physician’s Orders 5/7/2009, Eskalith – Lithium 300mg is to be taken 1 time daily Medication Administration Record & Physician’s Orders do not match.

Medication Administration Records did not contain the dosage of the medication given:
- Senna S – Ex-Lax – (1 to 2 tablets daily) 7/1, 2, 6, 7, 8, 16, 18, 23 & 30 (8:00 PM)

Individual #4
July 2009
As indicated by the Medication Administration Records the individual is to take Aspirin – Acetylsalicylic Acid 162 mg, 2 tablets (1 time daily). According to the Physician’s Orders, Aspirin – Acetylsalicylic Acid 162 mg is to be taken 1 time daily. Medication Administration Record & Physician’s Orders do not match. According to the MAR the individual is taking twice the prescribed dosage.

Individual #6
May 2009
Medication Administration Records contained missing entries. No documentation found indicating reason for missing entries:
- Tenormin – Atenolol 25mg (1 time daily) – Blank 5/2 (8:00 AM).
(i) Name of resident;
(ii) Date given;
(iii) Drug product name;
(iv) Dosage and form;
(v) Strength of drug;
(vi) Route of administration;
(vii) How often medication is to be taken;
(viii) Time taken and staff initials;
(ix) Dates when the medication is discontinued or changed;
(x) The name and initials of all staff administering medications.

Model Custodial Procedure Manual
D. Administration of Drugs

Unless otherwise stated by practitioner, patients will not be allowed to administer their own medications. Document the practitioner’s order authorizing the self-administration of medications.

All PRN (As needed) medications shall have complete detail instructions regarding the administering of the medication. This shall include:
- symptoms that indicate the use of the medication,
- exact dosage to be used, and
- the exact amount to be used in a 24 hour period.

Individual #7
June 2009
Medication Administration Records contained missing entries. No documentation found indicating reason for missing entries:
- Prilosec - Omeprazole dosage (1 time daily) – Blank 6/24 (4:00 PM)
- Coumadin – Warfarin 10mg (1 time daily) – Blank 6/24 (4:00 PM)
- Depakote – Valproic Acid dosage (2 times daily) – Blank 6/24 (4:00 PM)

Medication Administration Records did not contain the diagnosis for which the medication is prescribed:
- Valproic syrup 250mg/5 ml (2 times daily)

July 2009
Medication Administration Records did not contain the dosage of the medication which is to be given:
- Tegretol – Carbamazepine (2 times daily)
Tag # 1A09 Medication Delivery - PRN Medication

Scope and Severity Rating: E

Based on record review, the Agency failed to maintain PRN Medication Administration Records which contained all elements required by standard for 7 of 9 Individuals.

Individual #1
May 2009
No effectiveness was noted on the Medication Administration Record for the following PRN medication:
- Tylenol – Acetaminophen 325mg – PRN – 5/17 (given 2 times daily)

Individual #3
May 2009
Medication Administration Records did not contain the exact amount to be used in a 24 hour period:
- Tylenol – Acetaminophen 650mg (PRN)

Medication Administration Records did not contain the route of administration for the following medications:
- Tylenol – Acetaminophen 650mg (PRN)

No Signs/Symptoms were noted on the Medication Administration Record for the following PRN medication:
- Tylenol – Acetaminophen 650mg – PRN – 5/1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30 & 31 (given 1 time daily)

No Effectiveness was noted on the Medication Administration Record for the following PRN medication:
- Tylenol – Acetaminophen 650mg – PRN – 5/1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30 & 31 (given 1 time daily)
(f) For PRN medication, an explanation for the use of the PRN medication shall include observable signs/symptoms or circumstances in which the medication is to be used, and documentation of effectiveness of PRN medication administered.

(3) The Provider Agency shall also maintain a signature page that designates the full name that corresponds to each initial used to document administered or assisted delivery of each dose;

(4) MARs are not required for individuals participating in Independent Living who self-administer their own medications;

(5) Information from the prescribing pharmacy regarding medications shall be kept in the home and community inclusion service locations and shall include the expected desired outcomes of administering the medication, signs and symptoms of adverse events and interactions with other medications;

**NMAC 16.19.11.8 MINIMUM STANDARDS: A. MINIMUM STANDARDS FOR THE DISTRIBUTION, STORAGE, HANDLING AND RECORD KEEPING OF DRUGS:**

(d) The facility shall have a Medication Administration Record (MAR) documenting medication administered to residents, including over-the-counter medications. This documentation shall include:

(i) Name of resident;
(ii) Date given;
(iii) Drug product name;
(iv) Dosage and form;
(v) Strength of drug;
(vi) Route of administration;
(vii) How often medication is to be taken;
(viii) Time taken and staff initials;
(ix) Dates when the medication is discontinued

<table>
<thead>
<tr>
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<tbody>
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<td>Medication Administration Records did not contain the exact amount to be used in a 24 hour period:</td>
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<td>• Tylenol – Acetaminophen 650mg (PRN)</td>
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Medication Administration Records did not contain the route of administration for the following medications:

• Tylenol – Acetaminophen 650mg (PRN)

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No Effectiveness was noted on the Medication Administration Record for the following PRN medication:

• Tylenol – Acetaminophen 650mg – PRN – 6/1 - 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29 & 30 (given 1 time daily)

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Medication Administration Records did not contain the route of administration for the following medications:

• Tylenol – Acetaminophen 650mg (PRN)

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</tr>
</tbody>
</table>

DHI Quality Review Survey Report – ENMRSH, Inc. - Southeast Region – August 24 – 26, 2009

Report #: Q10.01.D1808.SE.001.FCD.01
or changed;

(x) The name and initials of all staff administering medications.

Model Custodial Procedure Manual

D. Administration of Drugs

Unless otherwise stated by practitioner, patients will not be allowed to administer their own medications. Document the practitioner’s order authorizing the self-administration of medications.

All PRN (As needed) medications shall have complete detail instructions regarding the administering of the medication. This shall include:

- symptoms that indicate the use of the medication,
- exact dosage to be used, and
- the exact amount to be used in a 24 hour period.

Department of Health

Developmental Disabilities Supports Division (DDSD) Medication Assessment and Delivery Policy - Eff. November 1, 2006

F. PRN Medication

3. Prior to self-administration, self-administration with physical assist or assisting with delivery of PRN medications, the direct support staff must contact the agency nurse to describe observed symptoms and thus assure that the PRN medication is being used according to instructions given by the ordering PCP. In cases of fever, respiratory distress (including coughing), severe pain, vomiting, diarrhea, change in responsiveness/level of consciousness, the nurse must strongly consider the need to conduct a face-to-face assessment to assure that the PRN does not mask a condition better treated by seeking medical attention. This does not apply to home based/family living settings where the provider is related by affinity or by consanguinity to the individual.

- Tylenol – Acetaminophen 650 mg – PRN – 7/1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30 & 31 (given 1 time daily)

No Effectiveness was noted on the Medication Administration Record for the following PRN medication:

- Tylenol – Acetaminophen 650mg – PRN – 7/1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30 & 31 (given 1 time daily)

Individual #5

May 2009

No Effectiveness was noted on the Medication Administration Record for the following PRN medication:

- Cheratussin AC/Robitussin AC – PRN – 5/17 (given 2 times daily)

Medication Administration Record did not contain the dosage of medication given:

- Cheratussin AC/Robitussin AC – PRN – 5/17 (given 2 times daily) & 5/20 & 24 (given 1 time daily)

Individual #6

May 2009

Medication Administration Records did not contain the circumstance for which the medication is to be used:

- Bentyl – Dicyclomine 20mg (PRN)

June 2009

Medication Administration Records did not contain the circumstance for which the medication is to be used:

- Bentyl – Dicyclomine 20mg (PRN)

July 2009

Medication Administration Records did not
4. The agency nurse shall review the utilization of PRN medications routinely. Frequent or escalating use of PRN medications must be reported to the PCP and discussed by the Interdisciplinary for changes to the overall support plan (see Section H of this policy).

H. Agency Nurse Monitoring

1. Regardless of the level of assistance with medication delivery that is required by the individual or the route through which the medication is delivered, the agency nurses must monitor the individual's response to the effects of their routine and PRN medications. The frequency and type of monitoring must be based on the nurse's assessment of the individual and consideration of the individual's diagnoses, health status, stability, utilization of PRN medications and level of support required by the individual's condition and the skill level and needs of the direct care staff. Nursing monitoring should be based on prudent nursing practice and should support the safety and independence of the individual in the community setting. The health care plan shall reflect the planned monitoring of the individual's response to medication.

| Department of Health Developmental Disabilities Supports Division (DDSD) - Procedure Title: Medication Assessment and Delivery Procedure
| Eff Date: November 1, 2006 |
| C. 3. Prior to delivery of the PRN, direct support staff must contact the agency nurse to describe observed symptoms and thus assure that the PRN is being used according to instructions given by the ordering PCP. In cases of fever, respiratory distress (including coughing), severe pain, vomiting, diarrhea, change in responsivenes/level of consciousness, the nurse must strongly consider the need to conduct a face-to-face assessment to assure that the PRN does not mask a condition |

| Individual #7 |
| May 2009 |
| Medication Administration Records did not contain the exact amount to be used in a 24 hour period: |
| • Reglan - Metoclopramide (PRN) |
| Medication Administration Records did not contain the circumstance for which the medication is to be given: |
| • Reglan - Metoclopramide (PRN) |
| • Tylenol – Acetaminophen 500mg (PRN) |
| Medication Administration Records did not contain the route of administration for the following medications: |
| • Reglan - Metoclopramide (PRN) |
| Medication Administration Records did not contain the strength of the medication which is to be given: |
| • Reglan – Metoclopramide (PRN) |
| • Albuterol Nebulizer (PRN) |

| No Signs/Symptoms were noted on the Medication Administration Record for the following PRN medication: |
| • Lortab Elixir – Acetaminophen & Hydrocodone 7.5/500 – 15 ml – PRN – 5/8 (given 1 time daily) |

| No Effectiveness was noted on the Medication Administration Record for the following PRN medication: |
| • Antipyrine – Benzocaine/Auralgan ear drops – PRN – 5/9 (given 1 time daily) |
better treated by seeking medical attention. (References: Psychotropic Medication Use Policy, Section D, page 5 Use of PRN Psychotropic Medications; and, Human Rights Committee Requirements Policy, Section B, page 4 Interventions Requiring Review and Approval – Use of PRN Medications).

4. Document on the MAR each time a PRN medication is used and describe its effect on the individual (e.g., temperature down, vomiting lessened, anxiety increased, the condition is the same, improved, or worsened, etc.).

<table>
<thead>
<tr>
<th>Date</th>
<th>No Effectiveness was noted on the Medication Administration Record for the following PRN medication:</th>
</tr>
</thead>
<tbody>
<tr>
<td>June 2009</td>
<td>Lortab Elixir – Acetaminophen &amp; Hydrocodone – PRN – 6/4 &amp; 5 (given 1 time daily)</td>
</tr>
<tr>
<td>July 2009</td>
<td>Tylenol – Acetaminophen – PRN – 7/21 (given 1 time daily)</td>
</tr>
<tr>
<td>Individual #8</td>
<td>No Effectiveness was noted on the Medication Administration Record for the following PRN medication:</td>
</tr>
<tr>
<td>May 2009</td>
<td>Tylenol – Acetaminophen 325mg – PRN – 5/6, 9, 12, 17, 19, 24, 26 &amp; 27 (given 1 time daily)</td>
</tr>
<tr>
<td>June 2009</td>
<td>Tylenol – Acetaminophen 325mg – PRN – 6/4, 9, 10, 14, 18, 27 &amp; 30 (given 1 time daily); 6/1, 2 &amp; 3 (given 2 times daily)</td>
</tr>
<tr>
<td>July 2009</td>
<td>Tylenol – Acetaminophen 325mg – PRN – 7/11, 12, 16 &amp; 17 (given 1 time daily)</td>
</tr>
<tr>
<td></td>
<td>Tylenol – Acetaminophen 325mg – PRN – 7/11, 12, 15, 16 &amp; 17 (given 1 time daily)</td>
</tr>
<tr>
<td>Individual #9</td>
<td>May 2009</td>
</tr>
<tr>
<td>----------------------</td>
<td>---------------------------------------------------------------------------</td>
</tr>
<tr>
<td></td>
<td>No Effectiveness was noted on the Medication Administration Record for the following PRN medication:</td>
</tr>
<tr>
<td></td>
<td>• Ibuprofen 600mg – PRN – 5/26 (given 1 time daily)</td>
</tr>
<tr>
<td></td>
<td>June 2009</td>
</tr>
<tr>
<td></td>
<td>No Effectiveness was noted on the Medication Administration Record for the following PRN medication:</td>
</tr>
<tr>
<td></td>
<td>• Ibuprofen 600mg – PRN – 6/8, 11 &amp; 15 (given 1 time daily) &amp; 6/24 (given 2 times daily)</td>
</tr>
</tbody>
</table>
### Tag # 1A27 (CoP) Duty to Report - IR’s Filed During On-Site and/or IR’s Not Reported by Provider

<table>
<thead>
<tr>
<th>Tag # 1A27 (CoP) Duty to Report - IR’s Filed During On-Site and/or IR’s Not Reported by Provider</th>
<th>Scope and Severity Rating: E</th>
</tr>
</thead>
</table>

#### 7.1.13.9 INCIDENT MANAGEMENT SYSTEM REPORTING REQUIREMENTS FOR COMMUNITY BASED SERVICE PROVIDERS: A. Duty To Report:
(1) All community based service providers shall immediately report abuse, neglect or misappropriation of property to the adult protective services division.
(2) All community based service providers shall report to the division within twenty four (24) hours: abuse, neglect, or misappropriation of property, unexpected and natural/expected deaths; and other reportable incidents to include:
- **a** an environmental hazardous condition, which creates an immediate threat to life or health; or
- **b** admission to a hospital or psychiatric facility or the provision of emergency services that results in medical care which is unanticipated or unscheduled for the consumer and which would not routinely be provided by a community based service provider.
(3) All community based service providers shall ensure that the reporter with direct knowledge of an incident has immediate access to the division incident report form to allow the reporter to respond to, report, and document incidents in a timely and accurate manner.

#### B. Notification:
(1) Incident Reporting: Any consumer, employee, family member or legal guardian may report an incident independently or through the community based service provider to the division by telephone call, written correspondence or other forms of communication utilizing the division’s incident report form. The incident report form and instructions for the completion and filing are available at the division's website: [http://dhi.health.state.nm.us/elibrary/ironline/ir.php](http://dhi.health.state.nm.us/elibrary/ironline/ir.php)

Based on record review, the Agency failed to report suspected abuse, neglect, or misappropriation of property, unexpected and natural/expected deaths; or other reportable incidents to the Division of Health Improvement for 2 of 8 Individuals. The following internal incidents were reported as a result of the on-site survey:

**Individual #1**
- Incident date May 2009 – July 2009 (Times vary). Medication Administration Records indicated the individual’s medications were not available, during the following dates: Aspirin - Acetylsalicylic Acid 325mg 5/5 – 19; Ditropan – Oxybutynin 5mg 6/4 – 6; Cogentin - Benzotropine 1mg 6/7 – 9; Seasonique–Levonorgestrel/Ethinyl Estradi 6/15 – 17; Lo-Ovral - Cryselle 28 7/14 – 18; Verelan - Verapamil 180mg; 7/14 – 16; Vitamin C 7/7 – 18; Multi Vitamin 7/7 – 18 & Os-Cal with Vitamin D 7/13 & 14. Incident report was filed on 8/26/2009 by the Agency.

**Individual #2**
- Incident date July 2009 (Times vary). Medication Administration Records indicated the individual’s medications were not given as prescribed, during the following dates: 7/1 - 13/2009 Eskalith – Lithium 300mg (2 times daily). According to the Physician’s Orders (5/7/2009), Eskalith – Lithium 300mg is to be taken 1 time daily. Incident report was filed on 8/26/2009 by the Agency.
or may be obtained from the department by calling the toll free number.

(2) Division Incident Report Form and Notification by Community Based Service Providers: The community based service provider shall report incidents utilizing the division’s incident report form consistent with the requirements of the division’s incident management system guide. The community based service provider shall ensure all incident report forms alleging abuse, neglect or misappropriation of consumer property submitted by a reporter with direct knowledge of an incident are completed on the division’s incident report form and received by the division within twenty-four (24) hours of an incident or allegation of an incident or the next business day if the incident occurs on a weekend or a holiday. The community based service provider shall ensure that the reporter with the most direct knowledge of the incident prepares the incident report form.

<table>
<thead>
<tr>
<th>Tag # 6L13 (CoP) - CL Healthcare Reqs.</th>
<th>Scope and Severity Rating: E</th>
</tr>
</thead>
<tbody>
<tr>
<td>CHAPTER 6. VI. GENERAL REQUIREMENTS FOR COMMUNITY LIVING</td>
<td></td>
</tr>
<tr>
<td>G. Health Care Requirements for Community Living Services.</td>
<td></td>
</tr>
<tr>
<td>(1) The Community Living Service providers shall ensure completion of a HAT for each individual receiving this service. The HAT shall be completed 2 weeks prior to the annual ISP meeting and submitted to the Case Manager and all other IDT Members. A revised HAT is required to also be submitted whenever the individual’s health status changes significantly. For individuals who are newly allocated to the DD Waiver program, the HAT may be completed within 2 weeks following the initial ISP meeting and submitted with any strategies and support plans indicated in the ISP, or within 72 hours following admission into direct services, whichever comes first.</td>
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<tr>
<td>(2) Each individual will have a Health Care Coordinator, designated by the IDT. When the individual’s HAT score is 4, 5 or 6 the Health Care Coordinator shall be an IDT member, other than the individual. The Health Care Coordinator shall oversee and monitor health care services for the individual in accordance with these standards. In circumstances where no IDT member voluntarily accepts designation as the health care coordinator, the community living provider shall assign a staff member to this role.</td>
<td></td>
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<tr>
<td>(3) For each individual receiving Community Living Services, the provider agency shall ensure and document the following:</td>
<td></td>
</tr>
<tr>
<td>a) Provision of health care oversight consistent with these Standards as detailed in Chapter One section III E: Healthcare Documentation by Nurses for Community Living Services, Community Inclusion Services and Private Duty Nursing Services.</td>
<td></td>
</tr>
<tr>
<td>b) That each individual with a score of 4, 5, or 6</td>
<td>Based on record review, the Agency failed to provide documentation of annual physical examinations and/or other examinations as specified by a licensed physician for 2 of 8 individuals receiving Community Living Services.</td>
</tr>
<tr>
<td>• Blood Pressure</td>
<td></td>
</tr>
<tr>
<td>° As indicated by Doctor’s Order 2/18/09, blood pressure should be taken 2 times weekly. None found 2/2009 - 7/2009 (#1)</td>
<td></td>
</tr>
<tr>
<td>• Blood Glucose</td>
<td></td>
</tr>
<tr>
<td>° As indicated by the Medication Administration Records May, June &amp; July 2009, blood glucose checks should be 3 times daily before meals and at bedtime. May 2009 through July 2009 Medication Administration Records document blood glucose checks done 3 times daily (8:00 AM, 12:00 PM &amp; 3:00 PM). Record does not indicate that blood glucose is being checked at “bedtime.” (#7)</td>
<td></td>
</tr>
</tbody>
</table>
on the HAT, has a Health Care Plan developed by a licensed nurse.
(c) That an individual with chronic condition(s) with the potential to exacerbate into a life threatening condition, has Crisis Prevention/Intervention Plan(s) developed by a licensed nurse or other appropriate professional for each such condition.
(4) That an average of 3 hours of documented nutritional counseling is available annually, if recommended by the IDT.
(5) That the physical property and grounds are free of hazards to the individual’s health and safety.
(6) In addition, for each individual receiving Supported Living or Family Living Services, the provider shall verify and document the following:
(a) The individual has a primary licensed physician;
(b) The individual receives an annual physical examination and other examinations as specified by a licensed physician;
(c) The individual receives annual dental check-ups and other check-ups as specified by a licensed dentist;
(d) The individual receives eye examinations as specified by a licensed optometrist or ophthalmologist; and
(e) Agency activities that occur as follow-up to medical appointments (e.g. treatment, visits to specialists, changes in medication or daily routine).