To: The Safety Division
   Pharmaceutical and Food Safety Bureau
   Ministry of Health, Labour and Welfare

Document of March 24, 2006
"Implementation of Bar Code Labeling of Ethical Drugs"

We thank you for the opportunity to review and comment on the above referenced
document. The global Healthcare User Group, GS1 HUG™ (www.gs1.org/hug),
comprises representation from major global pharmaceutical and medical device
manufacturers (including Japanese manufacturers), wholesalers, hospitals,
regulatory bodies and trade associations. The GS1 HUG™ is striving for global
standards for automatic product identification and is currently working with a number
of regulatory bodies.

The GS1 HUG™ Leadership team has reviewed this document in detail, together
with other GS1 HUG members and we would like to provide our comments, which
are listed in order of importance, some of which are recommendations for your
consideration and some of which require further clarification. We are available to
openly discuss these comments should you require clarification or additional
information.

Ref. 2 Numbering of product codes and JAN codes

The GS1 HUG is concerned about the requirements for packaging level indicators.
In the proposal, definitions are assigned to indicators 0, 1 and 2. The GS1
standards specify such indicators must be unique and do not have intelligence as
this limits flexibility for alternate package configurations. We strongly recommend
that the rule clarifies that packaging level indicators are not pre-assigned. The
manufacturer determines the packaging level indicator and ensures that each is
unique. GS1 HUG suggests that the table under section 3 (Changes of JAN codes)
be updated to incorporate this approach for JAN code changes.

Ref. 4 Bar code symbol system

The GS1 HUG suggests that any of the approved open GS1 standard symbologies
(including RSS, Data Matrix etc.) should be accepted. The market will drive the final
selection from the approved standards.

Ref. 5 Order for indicating data elements and application identifiers

According to GS1 standards, Application Identifier AI (30) is used for a variable
quantity, not a fixed quantity. AI (37) is to be used for a fixed quantity. The GS1
HUG strongly recommends that AI(30) and AI(37) are used as intended by GS1 standards. The case count field does not aid in preventing dispensing errors and therefore should not be within the scope of this proposed rule.

Ref. 7 (1) Others

This statement indicates two bar codes are required. The GS1 HUG recommends that only a single bar code is printed on any package unit.

Ref. 6 Timing for implementation of the New Bar Code Labeling (1)

It is unclear what products are classified as “specific biological products”. Is there a link to a database that can be shared? Is there logic to how specific biological products are identified?

Ref. ‘Ethical Drugs’

The wording ‘ethical drugs’ is used several times in the document. The GS1 HUG understanding is that an ethical drug is intended for the hospital market only? Clarification of the terminology should be considered.

Ref.1. (1) Formulation package unit

Clarification is needed around definition of the ‘formulation package unit’. Kits or combination packs may contain a vial of active substance and a vial of liquid for dilution, packaged together. What is defined as the smallest unit of package? The GS1 HUG assumes that the “unit of use” package is considered the formulation package unit.

The GS1 HUG will give serious consideration to suggest that the QR code should be included by GS1 as a future open standard.

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In the name of the GS1 HUG™ Leadership Team